

=> fil reg
FILE 'REGISTRY' ENTERED AT 14:57:15 ON 07 JUL 2006

=> d his ful

FILE 'HCAPLUS' ENTERED AT 13:59:35 ON 07 JUL 2006
L1 1 SEA ABB=ON US20050245755/PN
SEL RN

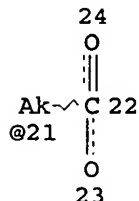
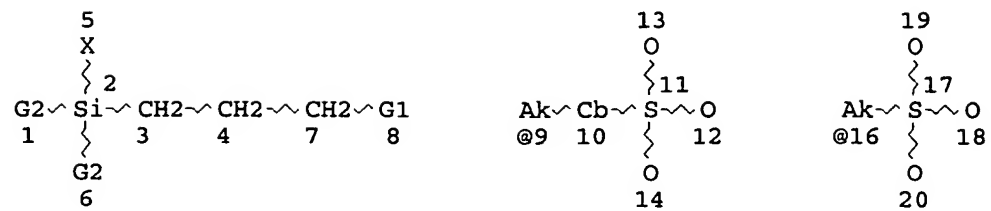
FILE 'REGISTRY' ENTERED AT 13:59:49 ON 07 JUL 2006
L2 7 SEA ABB=ON (10605-40-0/BI OR 1066-35-9/BI OR 107-05-1/
BI OR 12034-39-8/BI OR 13508-63-9/BI OR 298689-48-2/BI
OR 64-17-5/BI)

FILE 'REGISTRY' ENTERED AT 14:20:06 ON 07 JUL 2006
L3 STR
L4 0 SEA SSS SAM L3
D QUE STAT L4
L5 STR L3
L6 0 SEA SSS SAM L5
D QUE STAT
L7 37 SEA SSS FUL L5
L8 1 SEA ABB=ON L7 AND L2
SAV L7 NWA685/A

FILE 'HCAPLUS' ENTERED AT 14:44:19 ON 07 JUL 2006
L9 185 SEA ABB=ON L7
L10 47 SEA ABB=ON L9 AND (ALKANOL? OR ALCOHOL? OR ?ANOL?)
L11 1 SEA ABB=ON L10 AND L1
L12 2 SEA ABB=ON L10 AND LEATHER?/SC, SX
L13 43 SEA ABB=ON L10 AND (PROCESS? OR MAKING? OR SYNTHES?
OR PRODUC? OR PREP?)
L14 43 SEA ABB=ON (L11 OR L12 OR L13)
L15 37 SEA ABB=ON L14 AND (1840-2002)/PRY, AY, PY

FILE 'CASREACT' ENTERED AT 14:53:01 ON 07 JUL 2006
L16 STR L5
L17 0 SEA SSS SAM L16 (0 REACTIONS)
L18 8 SEA SSS FUL L16 (22 REACTIONS)
SAV L18 NWA685A/A

=> d que l15
L1 1 SEA FILE=HCAPLUS ABB=ON US20050245755/PN
L5 STR



VAR G1=X/9/16/21

VAR G2=AK/CB

NODE ATTRIBUTES:

DEFAULT MLEVEL IS ATOM

DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:

RING(S) ARE ISOLATED OR EMBEDDED

NUMBER OF NODES IS 23

STEREO ATTRIBUTES: NONE

L7 37 SEA FILE=REGISTRY SSS FUL L5
 L9 185 SEA FILE=HCAPLUS ABB=ON L7
 L10 47 SEA FILE=HCAPLUS ABB=ON L9 AND (ALKANOL? OR ALCOHOL?
 OR ?ANOL?)
 L11 1 SEA FILE=HCAPLUS ABB=ON L10 AND L1
 L12 2 SEA FILE=HCAPLUS ABB=ON L10 AND LEATHER?/SC, SX
 L13 43 SEA FILE=HCAPLUS ABB=ON L10 AND (PROCESS? OR MAKING?
 OR SYNTHES? OR PRODUC? OR PREP?)
 L14 43 SEA FILE=HCAPLUS ABB=ON (L11 OR L12 OR L13)
 L15 37 SEA FILE=HCAPLUS ABB=ON L14 AND (1840-2002)/PRY, AY, PY

=> fil hcap

FILE 'HCAPLUS' ENTERED AT 14:57:30 ON 07 JUL 2006

=> d l15 1-37 ibib abs hitstr hitind

L15 ANSWER 1 OF 37 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2004:293419 HCAPLUS

DOCUMENT NUMBER: 140:304720

TITLE: Production of silsesquioxane
derivative having functional group and
silsesquioxane derivativeINVENTOR(S): Yoshida, Kazuhiro; Ito, Kenya; Oikawa, Hisao;
Yamahiro, Mikio; Morimoto, Yoshitaka; Ohguma,
Koji; Watanabe, Kenichi; Ootake, Nobumasa

PATENT ASSIGNEE(S): Chisso Corporation, Japan

SOURCE: U.S. Pat. Appl. Publ., 24 pp.

DOCUMENT TYPE: CODEN: USXXCO
 LANGUAGE: Patent
 FAMILY ACC. NUM. COUNT: English 1
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 2004068074	A1	20040408	US 2003-661536	2003 0915
US 7053167	B2	20060530	<--	
JP 2005015738	A2	20050120	JP 2003-190847	2003 0703
US 2006089504	A1	20060427	US 2005-294364	2005 1206
PRIORITY APPLN. INFO.:			JP 2002-268716	A 2002 0913
			JP 2003-123678	A 2003 0428
			US 2003-661536	A1 2003 0915

OTHER SOURCE(S) : MARPAT 140:304720
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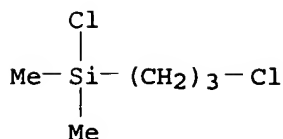
* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT
 *

AB A conventional silsesquioxane derivative has the problems that the functional groups are restricted and the chemical structure is not readily controlled, and that it is expensive. This is a process for producing a silsesquioxane derivative at a high yield by a simple low cost process. The silsesquioxane derivative represented by II, is characterized by using a Si compound salt intermediate material represented by I. Where in I and II, R = H, alkyl, aryl and arylalkyl; M = monovalent alkaline metal atom; ≥ 1 of Y = $-SiZR1R2$, and the remainder of Y = H; R1 and R2 = R; and Z = a functional group.

IT 10605-40-0, Chloro(3-chloropropyl)dimethylsilane
 (reaction with silsesquioxane salt; production of cage silsesquioxane derivative having functional group making use of salt intermediate)

RN 10605-40-0 HCAPLUS

CN Silane, chloro(3-chloropropyl)dimethyl- (6CI, 7CI, 8CI, 9CI) (CA INDEX NAME)



IC ICM C08G077-00
 INCL 528010000
 CC 37-6 (Plastics Manufacture and Processing)
 Section cross-reference(s): 29
 IT Silsesquioxanes
 (production of cage silsesquioxane derivative having
 functional group making use of salt intermediate)
 IT 429-60-7, Trimethoxy-3,3,3-trifluoropropylsilane 2996-92-1,
 Phenyltrimethoxysilane 18395-30-7, Isobutyltrimethoxysilane
 51851-37-7, Tridecafluoro-1,1,2,2-tetrahydrooctyltrithoxysilane
 143487-47-2, Cyclopentyltrimethoxysilane
 (condensation; production of cage silsesquioxane derivative
 having functional group making use of salt
 intermediate)
 IT 656800-15-6P
 (intermediate; production of cage silsesquioxane derivative
 having functional group making use of salt
 intermediate)
 IT 465499-97-2P 656800-11-2P 656800-14-5P
 (intermediate; production of cage silsesquioxane derivative
 having functional group making use of salt
 intermediate)
 IT 676229-30-4P 676229-37-1P 676616-38-9P 676616-39-0P
 676616-40-3P 676616-41-4P 676616-42-5P 676616-43-6P
 676616-44-7P 676616-45-8P 676616-46-9P 676616-47-0P
 676616-48-1P 676616-49-2P
 (production of cage silsesquioxane derivative having
 functional group making use of salt intermediate)
 IT 476635-00-4P
 (production of cage silsesquioxane derivative having
 functional group making use of salt intermediate)
 IT 106-92-3, Allyl glycidyl ether 111-45-5, 2-
 Allyloxyethanol 7539-12-0, Allylsuccinic anhydride
 13752-97-1 23523-56-0
 (reaction with H silsesquioxane; production of cage
 silsesquioxane derivative having functional group making
 use of salt intermediate)
 IT 1066-35-9, Chlorodimethylsilane 1481-41-0, Chlorodimethyl(3,3,3-
 trifluoropropyl)silane 1631-82-9, Chloromethylphenylsilane
 2227-29-4, Chlorodiisopropylsilane 10605-40-0,
 Chloro(3-chloropropyl)dimethylsilane 24636-31-5
 (reaction with silsesquioxane salt; production of cage
 silsesquioxane derivative having functional group making
 use of salt intermediate)

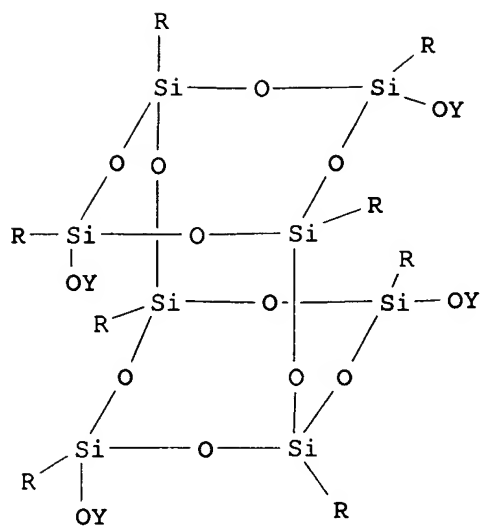
REFERENCE COUNT: 12 THERE ARE 12 CITED REFERENCES AVAILABLE
 FOR THIS RECORD. ALL CITATIONS AVAILABLE
 IN THE RE FORMAT

L15 ANSWER 2 OF 37 HCAPLUS COPYRIGHT 2006 ACS on STN
 ACCESSION NUMBER: 2004:252523 HCAPLUS
 DOCUMENT NUMBER: 140:272044

TITLE: Silsesquioxane derivative and process
 for producing the same
 INVENTOR(S): Yoshida, Kazuhiro; Morimoto, Yoshitaka;
 Watanabe, Kenichi; Ootake, Nobumasa
 PATENT ASSIGNEE(S): Chisso Corporation, Japan
 SOURCE: PCT Int. Appl., 56 pp.
 CODEN: PIXXD2
 DOCUMENT TYPE: Patent
 LANGUAGE: Japanese
 FAMILY ACC. NUM. COUNT: 1
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
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WO 2004024741	A1	20040325	WO 2003-JP11277	2003 0903
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RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG				
AU 2003261916	A1	20040430	AU 2003-261916	2003 0903
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US 2006052623	A1	20060309	US 2005-527751	2005 1019
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PRIORITY APPLN. INFO.:		JP 2002-268717	A	2002 0913
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		WO 2003-JP11277	W	2003 0903

OTHER SOURCE(S): MARPAT 140:272044
GI

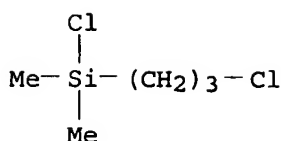


AB The present invention relates to (i) a silsesquioxane derivative I (PSQ derivative) which is for use as an electronic material, optical material, electrooptic material, or catalyst support and (ii) a process for producing the derivative, wherein R = H, alkyl, aryl, or arylalkyl; ≥ 1 of Ys = $\text{SiR}_1\text{R}_2\text{Z}$ (the remainder of Ys = H); R₁, R₂ = independently H, alkyl, aryl, or arylalkyl; and Z = functional group or functional group-containing group. Conventional PSQ derivs. have a problem that they have poor compatibility with general organic polymers. The novel PSQ derivative has improved compatibility with general organic polymers. The process enables the derivative to be produced in a short time at low cost. It is utilizable as an additive for improving the flame retardancy, heat resistance, weatherability, light resistance, elec. insulating properties, surface properties, hardness, mech. strength, chemical resistance, and other properties of general organic polymers. Thus, phenyltrimethoxysilane 6.54, water 0.66, and sodium hydroxide 0:88 kg, and 2-Pr alc. 26.3 L were refluxed for 5 h to give a silsesquioxane derivative, 69 g of which was reacted with 80 g chlorodimethylsilane to give a hydrosilyl-containing silsesquioxane, 2.0 g the resulting hydrosilyl-containing silsesquioxane was reacted with 1.4 g allyl glycidyl ether to give a 2.5 g glycidyl group-containing silsesquioxane with number average mol. weight 1100 and weight average mol. weight 1170.

IT 10605-40-0, 3-Chloropropyldimethylchlorosilane
(preparation of silsesquioxane derivs.)

RN 10605-40-0 HCPLUS

CN Silane, chloro(3-chloropropyl)dimethyl- (6CI, 7CI, 8CI, 9CI) (CA INDEX NAME)



IC ICM C07F007-21
 CC 38-3 (Plastics Fabrication and Uses)
 ST silsesquioxane deriv prep; phenyltrimethoxysilane
 chlorodimethylsilane allyl glycidyl ether contg silsesquioxane
 prep
 IT Silsesquioxanes
 (preparation of silsesquioxane derivs.)
 IT 502925-52-2P
 (intermediate; preparation of silsesquioxane derivs.)
 IT 502925-53-3P
 (model compound; preparation of silsesquioxane derivs.)
 IT 674298-98-7P
 (model compound; preparation of silsesquioxane derivs.)
 IT 674298-99-8P 674299-00-4P 674299-01-5P 674299-02-6P
 674299-03-7P 674299-04-8P
 (preparation of silsesquioxane derivs.)
 IT 106-92-3, Allyl glycidyl ether 111-45-5, 2-
 Allyloxyethanol 2996-92-1, Phenyltrimethoxysilane
 10605-40-0, 3-Chloropropyltrimethylchlorosilane
 23523-56-0, 4-Pentenoic acid trimethylsilyl ester 24636-31-5
 70964-99-7
 (preparation of silsesquioxane derivs.)
 REFERENCE COUNT: 2 THERE ARE 2 CITED REFERENCES AVAILABLE
 FOR THIS RECORD. ALL CITATIONS AVAILABLE
 IN THE RE FORMAT

L15 ANSWER 3 OF 37 HCPLUS COPYRIGHT 2006 ACS on STN
 ACCESSION NUMBER: 2003:1007372 HCPLUS
 DOCUMENT NUMBER: 140:43774
 TITLE: Method for preparation of
 organodialkylalkoxysilane
 INVENTOR(S): Ramdani, Kamel; Vigin, Bernard
 PATENT ASSIGNEE(S): Rhodia Chimie, Fr.; Rhone Poulenc Chimie
 SOURCE: Fr. Demande, 30 pp.
 CODEN: FRXXBL
 DOCUMENT TYPE: Patent
 LANGUAGE: French
 FAMILY ACC. NUM. COUNT: 1
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
FR 2841245	A1	20031226	FR 2002-7713	2002 0621
FR 2841245	B1	20050218		<--
FR 2841244	A1	20031226	FR 2002-15114	2002 1202
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WO 2004000852	A1	20031231	WO 2003-FR1921	
				2003 0623
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RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG				
AU 2003253076	A1	20040106	AU 2003-253076	
				2003 0623
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EP 1515977	A1	20050323	EP 2003-760774	
				2003 0623
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CN 1671719	A	20050921	CN 2003-818014	
				2003 0623
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JP 2005530855	T2	20051013	JP 2004-530906	
				2003 0623
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EP 1637534	A1	20060322	EP 2005-26550	
				2003 0623
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US 2005245755	A1	20051103	US 2005-518685	
				2005 0623
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PRIORITY APPLN. INFO.:		FR 2002-7713	A	
				2002 0621
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		FR 2002-15114	A	
				2002 1202
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		EP 2003-760774	A3	
				2003 0623
WO 2003-FR1921				
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				2003

0623

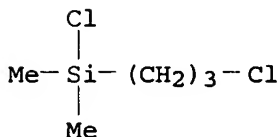
OTHER SOURCE(S) : CASREACT 140:43774; MARPAT 140:43774

AB The preparation of organodialkylalkoxysilane is carried out by reactive distillation of an ω -haloalkyldialkylhalosilane in the presence of an alkanol. The stage of reactive distillation is implemented in a column in the presence or absence of nonreactive solvent with the removal of HCl byproduct. The ω -haloalkyldialkylalkoxysilane thus obtained is particularly useful as starting material for preparation of organosilicon compds. containing sulfur and having general formula $R_1OSiR_2R_3(CH_2)3Sx(CH_2)3SiR_2R_3OR_1$ by reaction of sulfurization on an alkaline metal polysulfide.

IT 10605-40-0P, (3-Chloropropyl)dimethylchlorosilane
(method for preparation of organodialkylalkoxysilane)

RN 10605-40-0 HCPLUS

CN Silane, chloro(3-chloropropyl)dimethyl- (6CI, 7CI, 8CI, 9CI) (CA INDEX NAME)



IC ICM C07F007-18

CC 45-4 (Industrial Organic Chemicals, Leather, Fats, and Waxes)

Section cross-reference(s) : 37

ST haloalkyldialkylhalosilane distn organodialkylalkoxysilane prepns

IT 298689-48-2P, Bis[3-(dimethylethoxysilylpropyl)] tetrasulfide
(method for preparation of organodialkylalkoxysilane)

IT 64-17-5, Ethanol, reactions 107-05-1,
3-Chloropropylene 1066-35-9, Dimethylchlorosilane 12034-39-8,
Disodium tetrasulfide
(method for preparation of organodialkylalkoxysilane)

IT 10605-40-0P, (3-Chloropropyl)dimethylchlorosilane
13508-63-9P
(method for preparation of organodialkylalkoxysilane)

REFERENCE COUNT: 10 THERE ARE 10 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L15 ANSWER 4 OF 37 HCPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2003:837175 HCPLUS

DOCUMENT NUMBER: 139:308748

TITLE: Polysulfide siloxane applicable as vulcanizing agent and method for production thereof

INVENTOR(S): Belin, Laure; Blanchard, Christiane

PATENT ASSIGNEE(S): Societe de Technologie Michelin, Fr.; Michelin Recherche et Technique S.A.

SOURCE: PCT Int. Appl., 37 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE: French

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
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WO 2003087208	A1	20031023	WO 2003-EP3905	2003 0415
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RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG				
AU 2003227624	A1	20031027	AU 2003-227624	2003 0415
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EP 1499669	A1	20050126	EP 2003-725030	2003 0415
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EP 1499669	B1	20060111		
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JP 2005522515	T2	20050728	JP 2003-584160	2003 0415
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AT 315608	E	20060215	AT 2003-725030	2003 0415
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US 2005090680	A1	20050428	US 2004-945813	2004 0921
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PRIORITY APPLN. INFO.:			FR 2002-4964	A
				2002 0418
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			WO 2003-EP3905	W
				2003 0415

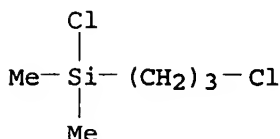
OTHER SOURCE(S): MARPAT 139:308748

AB Polysulfide disiloxanes in which the Si atoms are bridged by ZSxZ groups (Z = divalent group, x > 2) are manufactured for use as vulcanizing agents providing vulcanizates with improved heat stability.

IT 10605-40-0, 3-Chloropropylidemethylchlorosilane
(vulcanizing agent precursor; polysulfide disiloxanes

vulcanizing agents for vulcanizates with improved heat
resistance)

RN 10605-40-0 HCPLUS
CN Silane, chloro(3-chloropropyl)dimethyl- (6CI, 7CI, 8CI, 9CI) (CA
INDEX NAME)



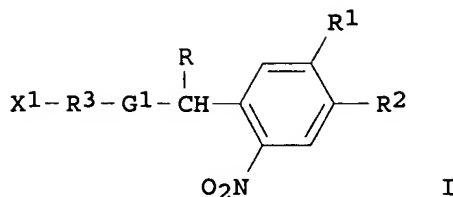
IC ICM C08K005-00
ICS C08J003-00; C08L009-00; C07F007-08
CC 39-10 (Synthetic Elastomers and Natural Rubber)
Section cross-reference(s): 29
IT 13508-63-9DP, cyclic polysulfide derivs. 174476-90-5DP, 3-
Chloropropyldimethylsilanol, cyclic polysulfide derivs.
(polysulfide disiloxanes vulcanizing agents for vulcanizates
with improved heat resistance)
IT 13508-63-9P 174476-90-5P, 3-Chloropropyldimethylsilanol
(vulcanizing agent precursor; polysulfide disiloxanes
vulcanizing agents for vulcanizates with improved heat
resistance)
IT 64-17-5, Ethanol, reactions 10605-40-0,
3-Chloropropyldimethylchlorosilane
(vulcanizing agent precursor; polysulfide disiloxanes
vulcanizing agents for vulcanizates with improved heat
resistance).

REFERENCE COUNT: 5 THERE ARE 5 CITED REFERENCES AVAILABLE
FOR THIS RECORD. ALL CITATIONS AVAILABLE
IN THE RE FORMAT

L15 ANSWER 5 OF 37 HCPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 2003:805778 HCPLUS
DOCUMENT NUMBER: 139:292355
TITLE: Preparation of silane coupling agent
INVENTOR(S): Yamaguchi, Kazuo; Ozaki, Atsushi
PATENT ASSIGNEE(S): Okamoto Chemical Industry Co., Ltd., Japan
SOURCE: Jpn. Kokai Tokkyo Koho, 19 pp.
CODEN: JKXXAF
DOCUMENT TYPE: Patent
LANGUAGE: Japanese
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
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JP 2003292496	A2	20031015	JP 2002-100926	2002 0403
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PRIORITY APPLN. INFO.:	JP 2002-100926			
	2002 0403			
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OTHER SOURCE(S):	MARPAT 139:292355			

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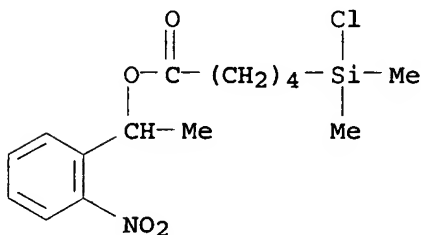


AB The patent relates to the preparation of nitrobenzyl alkoxy silyl derivs. I (G1 = O, COO; R1, R2 = H, methoxy etc.; R3 = methylene, alkylene etc.; X1 = trimethoxysilyl, triethoxysilyl; and R = H, alkyl etc.) as coupling agent useful for surface treatment of silicon wafer. Thus, 4,5-dimethoxy-2-nitrobenzyl 6-(trimethoxysilyl)hexyl ether prepared by reacting 5-hexenyl 4,5-dimethoxy-2-nitrobenzyl ether with trimethoxysilane was formulated in a composition comprising CST-70, CST-15, PSF2803, PSF2807, oil blue-613, and MEK to form a photo imaging solution which was coated on aluminum and gave pos. type picture after irradiation with mercury lamp at 365 nm.

IT 404353-10-2P
(preparation of nitrobenzyl alkoxy silyl coupling agent)

RN 404353-10-2 HCPLUS

CN Pentanoic acid, 5-(chlorodimethylsilyl)-, 1-(2-nitrophenyl)ethyl ester (9CI) (CA INDEX NAME)



IC ICM C07F007-18
ICS C08F008-42; C07F007-12
CC 29-6 (Organometallic and Organometalloidal Compounds)
Section cross-reference(s): 42, 74, 76
ST nitrobenzyl alkoxy silyl deriv coupling agent **prepn**
IT Diazotization
Hydrosilylation
(in preparation of nitrobenzyl alkoxy silyl coupling agent)
IT Coupling agents
(preparation of nitrobenzyl alkoxy silyl coupling agent)
IT Silanes
(preparation of nitrobenzyl alkoxy silyl coupling agent)
IT 3958-60-9
(in preparation of nitrobenzyl alkoxy silyl coupling agent)
IT 1016-58-6P 3205-25-2P 3718-21-6P 5385-87-5P 39716-58-0P,
4-Pentenoyl chloride 85834-41-9P 114119-94-7P 123830-38-6P

264258-90-4P 404353-09-9P 404353-12-4P 609355-38-6P
 609355-42-2P 609355-43-3P 609355-48-8P 609355-50-2P
 609355-51-3P
 (in preparation of nitrobenzyl alkoxy silyl coupling agent)
 IT 404353-13-5P
 (in preparation of nitrobenzyl alkoxy silyl coupling agent)
 IT 107-18-6, "Allyl alcohol", reactions 302-01-2,
 Hydrazine, reactions 591-80-0, 4-Pentenoic acid 614-21-1,
 "2-Nitroacetophenone" 712-97-0 821-41-0, "5-Hexene-1-ol"
 998-30-1, "Triethoxysilane" 1066-35-9, "Chlorodimethylsilane"
 2487-90-3, "Trimethoxysilane" 10025-78-2, "Trichlorosilane"
 13019-22-2, "9-Decen-1-ol" 20357-25-9 38460-95-6,
 10-Undecenoyl chloride 53169-26-9, "Chlorothionyl" 609355-44-4
 (in preparation of silane coupling agent)
 IT 609355-36-4P 609355-37-5P
 (in preparation of silane coupling agent)
 IT 404353-10-2P 404353-11-3P 404353-15-7P 404353-16-8P
 609355-39-7P 609355-40-0P 609355-41-1P 609355-45-5P
 609355-46-6P 609355-47-7P 609355-49-9P
 (preparation of nitrobenzyl alkoxy silyl coupling agent)

L15 ANSWER 6 OF 37 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2003:454333 HCAPLUS
 DOCUMENT NUMBER: 139:22334
 TITLE: Method for obtaining
 bis(monoorganoxysilylpropyl) polysulfides
 INVENTOR(S): Guennouni, Nathalie; Pevere, Virginie; Vigin, Bernard
 PATENT ASSIGNEE(S): Rhodia Chimie, Fr.
 SOURCE: PCT Int. Appl., 43 pp.
 CODEN: PIXXD2
 DOCUMENT TYPE: Patent
 LANGUAGE: French
 FAMILY ACC. NUM. COUNT: 1
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
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WO 2003048169	A1	20030612	WO 2002-FR4204	2002 1206

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W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA,
 CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI,
 GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG,
 KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK,
 MN, MW, MX, MZ, NO, NZ, OM, PH, PL, PT, RO, RU, SC, SD,
 SE, SG, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ,
 VC, VN, YU, ZA, ZM, ZW
 RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM,
 AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ,
 DE, DK, EE, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT,
 SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW,
 ML, MR, NE, SN, TD, TG

FR 2833264	A1	20030613	FR 2001-15768	2001 1206
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FR 2833264	B1	20050819		<--
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FR 2833265	A1	20030613	FR 2002-10145	
				2002 0809
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FR 2833265	B1	20060210		
AU 2002364429	A1	20030617	AU 2002-364429	
				2002 1206
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EP 1461344	A1	20040929	EP 2002-799785	
				2002 1206
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JP 2005511700	T2	20050428	JP 2003-549359	
				2002 1206
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EP 1621543	A1	20060201	EP 2005-21616	
				2002 1206
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R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, FI, CY, TR, BG, CZ, EE, SK				
PRIORITY APPLN. INFO.:				FR 2001-15768
				A
				2001 1206
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			FR 2002-10145	A
				2002 0809
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			EP 2002-799785	A3
				2002 1206
<--				
			WO 2002-FR4204	W
				2002 1206
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OTHER SOURCE(S): CASREACT 139:22334; MARPAT 139:22334

AB The invention concerns the preparation of bis(monoorganooxysilylpropyl) polysulfides $R_1OSiR_2R_3(CH_2)_3-Sx-(CH_2)_3SiR_2R_3OR_1$ (I, $R_1 = C_1-C_{15}$ alkyl, alkoxyalkyl; R_2 and $R_3 = C_1-C_6$ alkyl and/or phenyl; $1.5 \pm 1 \leq x \leq 5 \pm 0.1$). Said preparation is carried out by performing successively the following steps (a), (b) and (c): (a) hydrosilylation of the type: $R_2R_3HSi-Hal + CH_2:CH-CH_2-Hal \rightarrow Hal-R_2R_3Si-(CH_2)_3Hal$; (b) alcoholysis of the type: $Hal-R_2R_3Si-(CH_2)_3-Hal + R_1OH \rightarrow R_1O-R_2R_3Si-(CH_2)_3Hal$; (c) sulfidization of the type: $R_1O-R_2R_3Si-(CH_2)_3Hal + M_2Sx \rightarrow$ compound I; with Hal = halogen atom and M = alkali metal. Variations of the above reaction are also included in the invention. Thus, reaction of Me_2HSiCl with $CH_2:CHCH_2Cl$ in the presence of $[Ir(COD)Cl]_2$ (COD = 1,5-cyclooctadiene) as catalyst afforded $ClSiMe_2(CH_2)_3Cl$ (85% yield), which reacted with ethanol to give $EtOSiMe_2(CH_2)_3Cl$ (96% yield). Finally, reaction of the latter with Na_2S_4 afforded

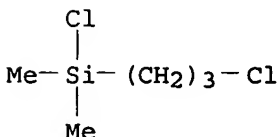
bis(monoorganooxysilylpropyl) tetrasulfide, EtOSiMe₂(CH₂)₃-S₄-(CH₂)₃SiMe₂OEt (87% yield).

IT 10605-40-0P

(intermediate; for preparation of
bis(monoethoxysilylpropyl) tetrasulfide)

RN 10605-40-0 HCPLUS

CN Silane, chloro(3-chloropropyl)dimethyl- (6CI, 7CI, 8CI, 9CI) (CA
INDEX NAME)



IC ICM C07F007-18

ICS C07F007-12; C07F007-14

CC 29-6 (Organometallic and Organometalloidal Compounds)

ST polysulfide monoorganooxysilylpropyl **prepn**;
hydrosilylation catalyst transition metal element compd allyl
halide alkoxy silane

IT Polysulfides

(preparation of bis(monoorganooxysilylpropyl) polysulfides)

IT Hydrosilylation catalysts

(transition metals and their compds. and complexes as catalysts
for hydrosilylation of allyl halide with alkoxy silanes, in
preparation of bis(monoorganooxysilylpropyl) polysulfides)

IT 12112-67-3, Chloro(1,5-cyclooctadiene)iridium dimer

(catalyst for hydrosilylation of allyl halide with
alkoxy silanes, in preparation of
bis(monoorganooxysilylpropyl) polysulfides)

IT 64-17-5, Ethanol, reactions 107-05-1, Allyl chloride
1066-35-9, Chlorodimethylsilane

(for preparation of bis(monoethoxysilylpropyl)
tetrasulfide)

IT 12034-39-8P, Disodium tetrasulfide
(for preparation of bis(monoethoxysilylpropyl)
tetrasulfide)

IT 10605-40-0P 13508-63-9P
(intermediate; for preparation of
bis(monoethoxysilylpropyl) tetrasulfide)

IT 298689-48-2P

(preparation of)

REFERENCE COUNT: 10 THERE ARE 10 CITED REFERENCES AVAILABLE
FOR THIS RECORD. ALL CITATIONS AVAILABLE
IN THE RE FORMAT

L15 ANSWER 7 OF 37 HCPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2002:863865 HCPLUS

DOCUMENT NUMBER: 139:69303

TITLE: Product subclass 42: γ -silyl
alkyl halides, alcohols, and esters
thereof

AUTHOR(S): Michael, J. P.; de Koning, C. B.

CORPORATE SOURCE: Molecular Science Institute, School of
Chemistry, University of Witwatersrand,
Johannesburg, 2050, S. Afr.

SOURCE: Science of Synthesis (2002), 4,

947-971

CODEN: SSCYJ9

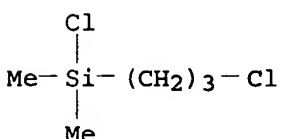
PUBLISHER: Georg Thieme Verlag
 DOCUMENT TYPE: Journal; General Review
 LANGUAGE: English

AB A review describes various methods for the synthesis of γ -silyl alkyl halides, alcs., and esters, and their applications. The methods described include the synthesis from silyl anions and functionalized three-carbon electrophiles; from silicon electrophiles and functionalized three-carbon nucleophiles; from α -silylated carbanions and epoxides; hydrosilylation of allylic compds.; coupling between vinylsilanes and aldehydes or ketones; addns. of allylsilanes; addition of β -silylated carbanions to aldehydes and ketones; and other miscellaneous methods.

IT 10605-40-0P
 (preparation of γ -silyl alkyl halides, alcs., and esters via hydrosilylation of allylic compound)

RN 10605-40-0 HCAPLUS

CN Silane, chloro(3-chloropropyl)dimethyl- (6CI, 7CI, 8CI, 9CI) (CA INDEX NAME)



CC 29-0 (Organometallic and Organometalloidal Compounds)

ST review gamma silyl alkyl halide alc ester synthesis; alkoxy silane retro Brook rearrangement review; allylic compd hydrosilylation review; vinyl silane aldehyde ketone coupling review; allyl silane hydrometalation oxidn review; radical addn allyl silane review; silyl enolate carbonyl compd condensation review

IT Rearrangement
 (Brook, retro; preparation of γ -silyl alkyl halides, alcs., and esters via retro-[1,4]-Brook rearrangement)

IT Addition reaction
 (allylboration; preparation of γ -silyl alkyl halides, alcs., and esters via allylboration-oxidation of allylsilanes)

IT Addition reaction
 (homolytic; preparation of γ -silyl alkyl halides, alcs., and esters via free-radical addition to allylsilanes)

IT Condensation reaction
 (preparation of γ -silyl alkyl halides, alcs., and esters via condensation of β -silyl enolates with carbonyl compound)

IT Coupling reaction
 (preparation of γ -silyl alkyl halides, alcs., and esters via coupling between vinylsilanes and aldehydes or ketones)

IT Hydrometalation
 Oxidation
 (preparation of γ -silyl alkyl halides, alcs., and esters via hydrometalation-oxidation of allylsilanes)

IT Hydrosilylation

(preparation of γ -silyl alkyl halides, alcs., and esters via hydrosilylation of allylic compound)

IT 98-60-2 763-13-3 1950-69-2 67957-50-0
(preparation of γ -silyl alkyl halides and alcs. from homoallylsilanes)

IT 18387-24-1P 67957-52-2P 210572-56-8P 210572-64-8P
(preparation of γ -silyl alkyl halides and alcs. from homoallylsilanes)

IT 18178-57-9 67957-95-3 75311-61-4
(preparation of γ -silyl alkyl halides via cleavage of silylcyclopropanes with acidic reagents)

IT 77508-38-4P 77508-44-2P 93297-66-6P 120347-42-4P
120347-49-1P 551939-66-3P
(preparation of γ -silyl alkyl halides via cleavage of silylcyclopropanes with acidic reagents)

IT 75-77-4, reactions 627-30-5 2203-35-2 14947-48-9,
1,3-Dithiane-2-ethanol 27607-77-8 53178-47-5
63823-55-2
(preparation of γ -silyl alkyl halides, alcs., and esters from silicon electrophiles and functionalized three-carbon electrophiles)

IT 2917-47-7P 52214-11-6P 94142-02-6P 124764-28-9P
129178-72-9P
(preparation of γ -silyl alkyl halides, alcs., and esters from silicon electrophiles and functionalized three-carbon electrophiles)

IT 94-41-7 503-30-0, Oxetane 930-68-7, 2-Cyclohexen-1-one
3839-31-4 32892-18-5 183729-68-2
(preparation of γ -silyl alkyl halides, alcs., and esters from silyl anions and functionalized three-carbon electrophiles)

IT 7452-98-4P 7452-99-5P 68469-62-5P 183729-69-3P
191916-53-7P 551939-22-1P
(preparation of γ -silyl alkyl halides, alcs., and esters from silyl anions and functionalized three-carbon electrophiles)

IT 75-21-8, Oxirane, reactions 75-56-9, reactions 185-70-6,
1-Oxaspiro[2.5]octane 286-20-4, 7-Oxabicyclo[4.1.0]heptane
762-72-1 1758-33-4 13683-41-5 17891-78-0 20780-53-4
123463-20-7 148259-36-3 159956-88-4
(preparation of γ -silyl alkyl halides, alcs., and esters from α -silylated carbanions and epoxides)

IT 86486-85-3P 103681-21-6P 144712-73-2P 159956-90-8P
174224-93-2P 551939-37-8P 551939-39-0P 551939-40-3P
551939-41-4P 551939-42-5P 551939-43-6P 551939-44-7P
551939-45-8P
(preparation of γ -silyl alkyl halides, alcs., and esters from α -silylated carbanions and epoxides)

IT 100-52-7, Benzaldehyde, reactions 123-19-3, 4-Heptanone
123-72-8, Butanal 700-58-3, Tricyclo[3.3.1.13,7]decanone
18156-67-7 104107-85-9 124853-60-7 164801-59-6
(preparation of γ -silyl alkyl halides, alcs., and esters via addition of β -silyl organometallic reagents to aldehydes or ketones)

IT 81372-27-2P 164801-60-9P 190380-95-1P 190380-97-3P
190381-00-1P 190381-02-3P 551939-62-9P
(preparation of γ -silyl alkyl halides, alcs., and esters via addition of β -silyl organometallic reagents to aldehydes or ketones)

IT 1113-12-8 10519-88-7 14579-08-9 24400-84-8 40934-71-2

184172-56-3 184172-57-4 551939-55-0
 (preparation of γ -silyl alkyl halides, alcs., and
 esters via allylboration-oxidation of allylsilanes)

IT 144930-15-4P 177594-75-1P 184172-60-9P 184172-61-0P
 184172-63-2P 184172-66-5P 184172-67-6P 184172-68-7P
 551939-56-1P
 (preparation of γ -silyl alkyl halides, alcs., and
 esters via allylboration-oxidation of allylsilanes)

IT 75-07-0, Acetaldehyde, reactions 18707-60-3 104085-59-8
 114431-81-1
 (preparation of γ -silyl alkyl halides, alcs., and
 esters via condensation of β -silyl enolates with carbonyl
 compound)

IT 104085-53-2P 104085-54-3P 104085-55-4P 104085-56-5P
 104113-42-0P 104113-43-1P 104113-44-2P 104113-46-4P
 146759-32-2P 146759-33-3P
 (preparation of γ -silyl alkyl halides, alcs., and
 esters via condensation of β -silyl enolates with carbonyl
 compound)

IT 67-64-1, 2-Propanone, reactions 78-93-3, 2-Butanone, reactions
 96-22-0, 3-Pentanone 98-86-2, reactions 108-94-1,
 Cyclohexanone, reactions 120-92-3, Cyclopentanone 754-05-2
 927-49-1, 6-Undecanone 2550-26-7 60484-87-9 66535-64-6
 101933-91-9 168282-42-6
 (preparation of γ -silyl alkyl halides, alcs., and
 esters via coupling between vinylsilanes and aldehydes or
 ketones)

IT 17888-67-4P 18410-35-0P 81372-29-4P 113386-95-1P
 125153-14-2P 158722-97-5P 158723-00-3P 168282-34-6P
 168282-41-5P 182947-90-6P 551939-53-8P
 (preparation of γ -silyl alkyl halides, alcs., and
 esters via coupling between vinylsilanes and aldehydes or
 ketones)

IT 3651-23-8 166970-54-3
 (preparation of γ -silyl alkyl halides, alcs., and
 esters via free-radical addition to allylsilanes)

IT 123331-54-4P 166970-55-4P
 (preparation of γ -silyl alkyl halides, alcs., and
 esters via free-radical addition to allylsilanes)

IT 29886-50-8 79753-70-1 97946-04-8 100312-73-0 106621-06-1
 146758-80-7
 (preparation of γ -silyl alkyl halides, alcs., and
 esters via hydrometalation-oxidation of allylsilanes)

IT 54040-91-4P 120196-04-5P 120196-05-6P 120196-19-2P
 120196-20-5P 146759-02-6P 146862-95-5P 551939-54-9P
 (preparation of γ -silyl alkyl halides, alcs., and
 esters via hydrometalation-oxidation of allylsilanes)

IT 75-36-5, Acetyl chloride 75-54-7 96-05-9 98-88-4, Benzoyl
 chloride 106-95-6, reactions 107-05-1 107-18-6,
 2-Propen-1-ol, reactions 563-47-3 563-52-0 591-87-7
 998-30-1 999-55-3 1066-35-9 1438-82-0 1631-70-5
 1631-84-1 3282-30-2 10025-78-2 14857-34-2 18146-00-4
 58210-84-7 138924-51-3 139016-94-7 139016-95-8 139016-97-0
 139017-00-8 551939-46-9
 (preparation of γ -silyl alkyl halides, alcs., and
 esters via hydrosilylation of allylic compound)

IT 1000-58-4P 1591-20-4P 1628-11-1P 2550-06-3P 3401-26-1P
 5089-70-3P 5290-24-4P 7787-93-1P 10605-40-0P
 13883-39-1P 18142-53-5P 18209-82-0P 18301-56-9P
 18387-98-9P 37611-45-3P 38595-89-0P 42496-16-2P

58210-63-2P 61676-44-6P 551939-47-0P 551939-48-1P
 551939-49-2P 551939-50-5P 551939-51-6P 551939-52-7P
 551939-67-4P 551939-68-5P
 (preparation of γ -silyl alkyl halides, alcs., and
 esters via hydrosilylation of allylic compound)
 IT 18171-15-8 18244-07-0 141412-20-6 190523-14-9 198646-26-3
 198646-27-4 198646-30-9 198646-31-0 551939-24-3
 551939-25-4 551939-28-7
 (preparation of γ -silyl alkyl halides, alcs., and
 esters via retro-[1,4]-Brook rearrangement)
 IT 18387-35-4P 141437-79-8P 150845-21-9P 150845-22-0P
 163123-02-2P 163123-13-5P 198646-19-4P 198646-20-7P
 198646-34-3P 551939-23-2P 551939-26-5P 551939-27-6P
 (preparation of γ -silyl alkyl halides, alcs., and
 esters via retro-[1,4]-Brook rearrangement)
 IT 78-79-5, reactions 110-62-3, Pentanal 995-45-9 2288-18-8
 87436-97-3
 (preparation of γ -silyl alkyl halides, alcs., and
 esters via silylmetalation of dienes followed by addition to
 carbonyl compound)
 IT 81906-08-3P 133447-98-0P 133447-99-1P 133448-00-7P
 133448-01-8P 133448-06-3P 187098-09-5P 187098-12-0P
 551939-65-2P
 (preparation of γ -silyl alkyl halides, alcs., and
 esters via silylmetalation of dienes followed by addition to
 carbonyl compound)

REFERENCE COUNT: 118 THERE ARE 118 CITED REFERENCES AVAILABLE
 FOR THIS RECORD. ALL CITATIONS AVAILABLE
 IN THE RE FORMAT

L15 ANSWER 8 OF 37 HCAPLUS COPYRIGHT 2006 ACS on STN
 ACCESSION NUMBER: 2002:658673 HCAPLUS
 DOCUMENT NUMBER: 137:165805
 TITLE: Low fluorescence nylon/glass composites for
 micro-analytical diagnostic applications
 INVENTOR(S): Andreoli, Rita; Amin, Murtaza; Meyering, Mark;
 Chesterson, Richard; Ostreicher, Eugene
 PATENT ASSIGNEE(S): Cuno Inc., USA
 SOURCE: U.S. Pat. Appl. Publ., 21 pp., Cont.-in-part
 of U.S. Provisional Ser. No. 224,141.
 CODEN: USXXCO
 DOCUMENT TYPE: Patent
 LANGUAGE: English
 FAMILY ACC. NUM. COUNT: 8
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 2002119559	A1	20020829	US 2001-899607	2001 0705 ---
US 6734012	B2	20040511		
US 2004157320	A1	20040812	US 2004-772645	2004 0205 ---
PRIORITY APPLN. INFO.:			US 2000-216229P	P 2000 ---

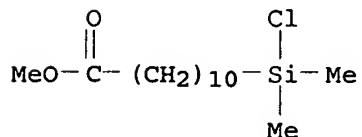
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AB An improved combination non-luminescent microporous membrane and solid support for use in micro-anal. diagnostic applications is disclosed. Specifically, a multi-cell non-luminescent substrate having a porous membrane formed by a phase inversion process effectively attached by covalent bonding through a surface treatment to a substrate that preps. the substrate to sufficiently, covalently bond to the non-luminescent microporous membrane formed by a phase inversion process such that the combination produced thereby is useful in microarray applications and wherein the porous non-luminescent nylon multi-cell substrate is covalently bonded to a solid base member, such as, for example, a glass or Mylar microscope slide, such that the combination produced thereby is useful in microarray applications. Apparatus and methods for fabricating the non-luminescent multi-cell substrate are also disclosed.

IT 53749-38-5, (10-Carbomethoxydecyl)dimethylchlorosilane (improved low fluorescence nylon/glass composites for micro-anal. diagnostic applications)

RN 53749-38-5 HCAPLUS

CN Undecanoic acid, 11-(chlorodimethylsilyl)-, methyl ester (9CI) (CA INDEX NAME)



IC ICM C12M001-34
ICS B05D003-00

INCL 435287200

CC 9-1 (Biochemical Methods)
Section cross-reference(s): 35, 47

IT 64-17-5, Ethanol, uses 67-56-1, Methanol, uses 112-57-2, Tetraethylenepentamine 7732-18-5, Water, uses (improved low fluorescence nylon/glass composites for micro-anal. diagnostic applications)

IT 64-18-6, Formic acid, reactions 106-89-8D, Epichlorohydrin, reaction products with polyamide-polyamines 919-30-2, 3-Aminopropyl triethoxysilane 1760-24-3, N-(2-Aminoethyl)-3-aminopropyl trimethoxysilane 2530-83-8, 3-Glycidoxypropyltrimethoxysilane 3388-04-3, 2-(3,4-Epoxy cyclohexyl)ethyltrimethoxysilane 53749-38-5, (10-Carbomethoxydecyl)dimethylchlorosilane

(improved low fluorescence nylon/glass composites for
micro-anal. diagnostic applications)

REFERENCE COUNT: 23 THERE ARE 23 CITED REFERENCES AVAILABLE
FOR THIS RECORD. ALL CITATIONS AVAILABLE
IN THE RE FORMAT

L15 ANSWER 9 OF 37 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 2002:293753 HCAPLUS
DOCUMENT NUMBER: 136:311081
TITLE: Rubber composition comprising, as coupling
agent, a polyfunctional organosilane
INVENTOR(S): Tardivat, Jean-Claude; Pagano, Salvatore
PATENT ASSIGNEE(S): Societe De Technologie Michelin, Fr.; Michelin
Recherche Et Technique S.A.
SOURCE: PCT Int. Appl., 41 pp.
CODEN: PIXXD2
DOCUMENT TYPE: Patent
LANGUAGE: French
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
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WO 2002031041	A1	20020418	WO 2001-EP11669	2001 1009
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W: AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CR, CU, CZ, DE, DK, DM, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW				
RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG				
CA 2425330	AA	20020418	CA 2001-2425330	2001 1009
<--				
AU 2002023607	A5	20020422	AU 2002-23607	2001 1009
<--				
EP 1326914	A1	20030716	EP 2001-986704	2001 1009
<--				
EP 1326914	B1	20060621		
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR				
BR 2001014614	A	20031223	BR 2001-14614	2001 1009
<--				
JP 2004511601	T2	20040415	JP 2002-534417	2001 1009

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 US 2004051210 A1 20040318 US 2003-411615
 2003
 0410

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 PRIORITY APPLN. INFO.: FR 2000-13255 A 2000
 2000
 1013
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 WO 2001-EP11669 W 2001
 2001
 1009
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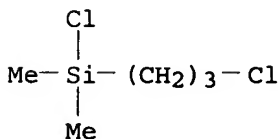
OTHER SOURCE(S): MARPAT 136:311081

AB The invention concerns an elastomeric composition based on at least a diene elastomer, an inorg. filler as reinforcing filler, a polyfunctional organosilane as coupling agent (inorg. filler/diene elastomer), bearing at least two functions designated X and Y, capable of being grafted on the elastomer by function X and on the inorg. filler by function Y. The invention is characterized in that said function Y is a hydroxysilyl function. Preferably, said organosilane is a hydroxysilane polysulfide, the diene elastomer is selected in the group consisting of polybutadiene, natural rubber, synthetic polyisoprene, butadiene copolymers, isoprene copolymers, and the reinforcing inorg. filler is a highly dispersible silicious filler. The compns. containing these silanes exhibit improved scorching properties and green-state processability. The invention also concerns tires or semi-finished products for tires, in particular running treads for tires comprising the inventive composition. A typical coupling agent was manufactured by reaction of ClSiMe₂(CH₂)₃Cl (I) with EtOH in the presence of NET₃ and reaction of the resulting EtOSiMe₂(CH₂)₃Cl with (1) MeOH/aqueous NaOH, (2) aqueous KH₂PO₄, and (3) ether or by reaction of I with NET₃, H₂O, ET₂O and reaction of the resulting HOSiMe₂(CH₂)₃Cl with Na₂Sx in the presence of H₂O, NaCl, PhMe, and a phase-transfer catalyst.

IT 10605-40-0, 3-Chloropropyldimethylchlorosilane
 (coupling agent precursor; rubber compns. containing hydroxysilyl polysulfide coupling agents for silica in tires)

RN 10605-40-0 HCAPLUS

CN Silane, chloro(3-chloropropyl)dimethyl- (6CI, 7CI, 8CI, 9CI) (CA INDEX NAME)



IC ICM C08K005-548
 ICS C08L021-00

CC 39-13 (Synthetic Elastomers and Natural Rubber)

IT 64-17-5, Ethanol, reactions 10605-40-0,
 3-Chloropropyldimethylchlorosilane
 (coupling agent precursor; rubber compns. containing hydroxysilyl polysulfide coupling agents for silica in tires)

IT 1344-08-7DP, Sodium polysulfide, reaction products with
 chloropropyldimethylhydroxysilane 174476-90-5DP, reaction

products with sodium polysulfide
 (rubber compns. containing hydroxysilyl polysulfide coupling agents
 for silica in tires)

REFERENCE COUNT: 4 THERE ARE 4 CITED REFERENCES AVAILABLE
 FOR THIS RECORD. ALL CITATIONS AVAILABLE
 IN THE RE FORMAT

L15 ANSWER 10 OF 37 HCAPLUS COPYRIGHT 2006 ACS on STN
 ACCESSION NUMBER: 2002:293661 HCAPLUS
 DOCUMENT NUMBER: 136:311046
 TITLE: Polyfunctional organosilanes for use as
 coupling agent for diene rubber
 Inventor(s): Tardivat, Jean-Claude; Belin, Laure;
 Blanchard, Christine
 PATENT ASSIGNEE(S): Societe De Technologie Michelin, Fr.; Michelin
 Recherche Et Technique S.A.
 SOURCE: PCT Int. Appl., 29 pp.
 CODEN: PIXXD2
 DOCUMENT TYPE: Patent
 LANGUAGE: French
 FAMILY ACC. NUM. COUNT: 1
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2002030939	A1	20020418	WO 2001-EP11668	2001 1009
<--				
W: AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CR, CU, CZ, DE, DK, DM, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW				
RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG				
CA 2425300	AA	20020418	CA 2001-2425300	2001 1009
<--				
AU 2002016955	A5	20020422	AU 2002-16955	2001 1009
<--				
EP 1326871	A1	20030716	EP 2001-986691	2001 1009
<--				
EP 1326871	B1	20060201		
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR				
BR 2001014616	A	20031223	BR 2001-14616	2001 1009
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JP 2004520272	T2	20040708	JP 2002-534324	

AT 316973	E	20060215	AT 2001-986691	2001 1009
RU 2272042	C2	20060320	RU 2003-113527	2001 1009
US 6774255	B1	20040810	US 2003-411616	2003 0410
PRIORITY APPLN. INFO.:			FR 2000-13254	A 2000 1013
			WO 2001-EP11668	W 2001 1009
<--				

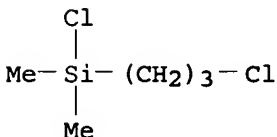
OTHER SOURCE(S): MARPAT 136:311046

AB The invention concerns monohydroxysilane polysulfide, $\text{HOSiR}_2(\text{R}'\text{S}_x\text{R}')\text{SiR}_2\text{OH}$, wherein: the radicals R, identical or different, are hydrocarbon groups comprising preferably 1 to 15 C atoms; the radicals R', identical or different, are divalent binding groups preferably comprising 1 to 18 C atoms; x is ≥ 2 . Said hydroxysilane is in particular a bis(propyldimethylsilanol) sulfide. The method for obtaining said hydroxysilane consists in subjecting to alcoholysis and/or hydrolysis a halogenated organosilane, followed by a sulfidizing step. For example, $\text{ClSiMe}_2(\text{CH}_2)_3\text{Cl}$ was either converted to $\text{EtOSiMe}_2(\text{CH}_2)_3\text{Cl}$ ($\text{EtOH/Et}_3\text{N}$) and then sequentially treated with (1) MeOH/NaOHaq , (2) $\text{KH}_2\text{PO}_4\text{aq}$ and (3) Et_2O to give $\text{HOSiMe}_2(\text{CH}_2)_3\text{Cl}$ or treated with $\text{Et}_3\text{N/H}_2\text{O/Et}_2\text{O}$ to give $\text{HOSiMe}_2(\text{CH}_2)_3\text{Cl}$, which was then reacted with $\text{Na}_2\text{S}_x/\text{H}_2\text{O/NaCl/toluene/Bu}_4\text{NBr}$ to give $\text{HOSiMe}_2(\text{CH}_2)_3\text{S}_x(\text{CH}_2)_3\text{SiMe}_2\text{OH}$ ($x = 2-6$ with average value ≈ 3.7). The invention also concerns the use of said hydroxysilane as coupling agent.

IT 10605-40-0, Chloro(3-chloropropyl)dimethylsilane
(for preparation of hydroxysilane polysulfides useful as coupling agents for diene elastomers)

RN 10605-40-0 HCAPLUS

CN Silane, chloro(3-chloropropyl)dimethyl- (6CI, 7CI, 8CI, 9CI) (CA INDEX NAME)



IC ICM C07F007-08

ICS C08K005-31; C08K005-548

CC 39-2 (Synthetic Elastomers and Natural Rubber)

Section cross-reference(s): 29

ST hydroxysilane polysulfide prep use coupling agent SBR;
 silanol polysulfide coupling agent butyl rubber
 IT Polysulfides
 Silanes
 (monohydroxysilane polysulfides; preparation for use as
 coupling agents for diene elastomers)
 IT Butyl rubber, properties
 (monohydroxysilane polysulfides; preparation for use as
 coupling agents for diene elastomers)
 IT Styrene-butadiene rubber, properties
 (monohydroxysilane polysulfides; preparation for use as
 coupling agents for diene elastomers)
 IT Coupling agents
 (preparation of hydroxysilane polysulfides for diene
 elastomers)
 IT 9010-85-9
 (butyl rubber, monohydroxysilane polysulfides; preparation
 for use as coupling agents for diene elastomers)
 IT 7631-86-9, Zeosil 1165MP, uses
 (for preparation of hydroxysilane polysulfides useful as
 coupling agents for diene elastomers)
 IT 10605-40-0, Chlоро(3-chloropropyl)dimethylsilane
 (for preparation of hydroxysilane polysulfides useful as
 coupling agents for diene elastomers)
 IT 13508-63-9P, (3-Chloropropyl)(ethoxy)dimethylsilane
 174476-90-5P, (3-Chloropropyl)dimethylsilanol
 (intermediate; for preparation of hydroxysilane
 polysulfides useful as coupling agents for diene elastomers)
 IT 411223-82-0P, Bis(3-(hydroxydimethylsilyl)propyl) disulfide
 411223-83-1P, Bis(3-(hydroxydimethylsilyl)propyl) trisulfide
 411223-84-2P, Bis(3-(hydroxydimethylsilyl)propyl) tetrasulfide
 411223-85-3P, Bis(3-(hydroxydimethylsilyl)propyl) pentasulfide
 411223-86-4P
 (preparation for use as coupling agents for diene
 elastomers)
 IT 9003-55-8
 (styrene-butadiene rubber, monohydroxysilane polysulfides;
 preparation for use as coupling agents for diene elastomers)

REFERENCE COUNT: 1 THERE ARE 1 CITED REFERENCES AVAILABLE
 FOR THIS RECORD. ALL CITATIONS AVAILABLE
 IN THE RE FORMAT

L15 ANSWER 11 OF 37 HCPLUS COPYRIGHT 2006 ACS on STN
 ACCESSION NUMBER: 2002:205086 HCPLUS
 DOCUMENT NUMBER: 136:247695
 TITLE: Nitrobenzyl group-containing chlorosilanes as
 coupling agents, and introduction of carboxy
 or hydroxy group to material surfaces using
 them
 INVENTOR(S): Yamaguchi, Kazuo; Futami, Tatsuhiro
 PATENT ASSIGNEE(S): Okamoto Kagaku Kogyo K. K., Japan
 SOURCE: Jpn. Kokai Tokkyo Koho, 14 pp.
 CODEN: JKXXAF
 DOCUMENT TYPE: Patent
 LANGUAGE: Japanese
 FAMILY ACC. NUM. COUNT: 1
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
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JP 2002080481 A2 20020319 JP 2000-2699042000
0906

PRIORITY APPLN. INFO.: JP 2000-269904

2000
0906

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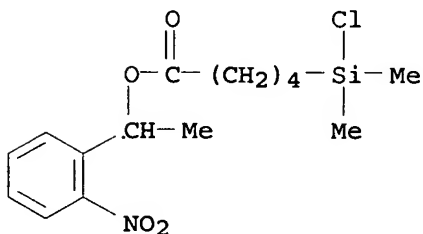
OTHER SOURCE(S): MARPAT 136:247695

AB The compds. have SiClMe₂, SiCl₂Me, or SiCl₃ at one terminal and (α -substituted) p-nitrobenzyl group at the other terminal. 1-(2-Nitrophenyl)ethyl 4-pentenoate (preparation given) was hydrosilylated by HSiCl₃ in the presence of H₂PtCl₆ to give 79% Cl₃Si(CH₂)₄CO₂CHMeC₆H₄NO₂-o (I). A Si wafer was treated with C₆H₆ solution of I and irradiated by UV to give a surface-modified wafer with high contact angle.

IT 404353-10-2P (preparation of nitrobenzyl group-containing silane coupling agents)

RN 404353-10-2 HCPLUS

CN Pentanoic acid, 5-(chlorodimethylsilyl)-, 1-(2-nitrophenyl)ethyl ester (9CI) (CA INDEX NAME)



IC ICM C07F007-08

ICS C07F007-08; C09C003-12

CC 29-6 (Organometallic and Organometalloidal Compounds)

IT Coupling agents (preparation of nitrobenzyl group-containing silane coupling agents)

IT 404353-10-2P 404353-11-3P 404353-15-7P 404353-16-8P (preparation of nitrobenzyl group-containing silane coupling agents)

IT 552-89-6, 2-Nitrobenzaldehyde 591-80-0, 4-Pentenoic acid 614-21-1, 2-Nitroacetophenone 821-41-0, 5-Hexen-1-ol 1066-35-9, Chlorodimethylsilane 3958-60-9, 2-Nitrobenzyl bromide 10025-78-2, Trichlorosilane 13019-22-2, 9-Decen-1-ol (preparation of nitrobenzyl group-containing silane coupling agents)

IT 3205-25-2P, 1-(2-Nitrophenyl)ethanol 3718-21-6P 5385-87-5P 39716-58-0P, 4-Pentenoic acid chloride 85834-41-9P 116271-34-2P 264258-90-4P 404353-09-9P 404353-12-4P 404353-13-5P 404353-14-6P (preparation of nitrobenzyl group-containing silane coupling agents)

IT 7440-21-3, Silicon, processes (wafer; preparation of nitrobenzyl group-containing silane coupling agents for)

L15 ANSWER 12 OF 37 HCAPLUS COPYRIGHT 2006 ACS on STN
 ACCESSION NUMBER: 2002:31474 HCAPLUS
 DOCUMENT NUMBER: 136:82254
 TITLE: Improved low fluorescence nylon/glass
 composites for micro-analytical diagnostic
 applications
 INVENTOR(S): Andreoli, Rita; Amin, Murtaza; Myering, Mark;
 Chesterton, Richard; Ostreicher, Eugene
 PATENT ASSIGNEE(S): Cuno, Inc., USA
 SOURCE: PCT Int. Appl., 43 pp.
 CODEN: PIXXD2
 DOCUMENT TYPE: Patent
 LANGUAGE: English
 FAMILY ACC. NUM. COUNT: 8
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
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WO 2002002585	A2	20020110	WO 2001-US21262	2001 0705
WO 2002002585	A3	20021003	-----	-----
W: AU, BR, JP RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR				
EP 1297340	A2	20030402	EP 2001-950882	2001 0705
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, FI, CY, TR			-----	-----
JP 2004502928	T2	20040129	JP 2002-507837	2001 0705
BR 2001006969	A	20040713	BR 2001-6969	2001 0705
AU 780441	B2	20050324	AU 2001-71835	2001 0705
PRIORITY APPLN. INFO.:			US 2000-216229P	P 2000 0705
			US 2000-216390P	P 2000 0706
			US 2000-224141P	P 2000 0810
			WO 2001-US21262	W 2001

0705

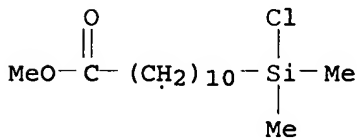
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AB An improved combination non-luminescent microporous membrane and solid support for use in micro-anal. diagnostic applications is disclosed. Specifically, a multi-cell non-luminescent substrate having a porous membrane formed by a phase inversion process effectively attached by covalent bonding through a surface treatment to a substrate that preps. the substrate to sufficiently, covalently bond to the non-luminescent microporous membrane formed by a phase inversion process such that the combination produced thereby is useful in microarray applications and wherein the porous non-luminescent nylon multi-cell substrate is covalently bonded to a solid base member, such as, for example, a glass or Mylar microscope slide, such that the combination produced thereby is useful in microarray applications. Apparatus and methods for fabricating the non-luminescent multi-cell substrate are also disclosed. A carbon black-impregnated nylon membrane was cast and then permanently attached to a glass slide to form a composite.

IT 53749-38-5, (10-Carbomethoxydecyl)dimethylchlorosilane (improved low fluorescence nylon/glass composites for micro-anal. diagnostic applications)

RN 53749-38-5 HCAPLUS

CN Undecanoic acid, 11-(chlorodimethylsilyl)-, methyl ester (9CI) (CA INDEX NAME)



IC ICM C07H021-00

CC 9-1 (Biochemical Methods)
Section cross-reference(s): 35, 47

IT 64-17-5, Ethanol, uses 67-56-1, Methanol, uses 112-57-2, Tetraethylenepentamine 7732-18-5, Water, uses (improved low fluorescence nylon/glass composites for micro-anal. diagnostic applications)

IT 64-18-6, Formic acid, reactions 106-89-8D, Epichlorohydrin, reaction products with polyamide-polyamines 919-30-2, 3-Aminopropyl triethoxysilane 1760-24-3, N-(2-Aminoethyl)-3-aminopropyl trimethoxysilane 2530-83-8, 3-Glycidoxypropyltrimethoxysilane 3388-04-3, 2-(3,4-Epoxy cyclohexyl)ethyltrimethoxysilane 53749-38-5, (10-Carbomethoxydecyl)dimethylchlorosilane (improved low fluorescence nylon/glass composites for micro-anal. diagnostic applications)

L15 ANSWER 13 OF 37 HCAPLUS COPYRIGHT 2006 ACS on STN
 ACCESSION NUMBER: 2001:832173 HCAPLUS
 DOCUMENT NUMBER: 136:111860
 TITLE: Immobilization of difunctional building blocks on hydroxysuccinimide activated silica: versatile in situ preparation of chiral stationary phases
 AUTHOR(S): Kosjek, Birgit; Uray, Georg
 CORPORATE SOURCE: Institut fur Chemie, Karl-Franzens Universitat

SOURCE: Graz, Graz, 8010, Austria
 Chirality (2001), 13(10), 657-667
 CODEN: CHRLEP; ISSN: 0899-0042

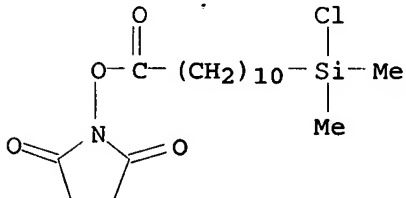
PUBLISHER: Wiley-Liss, Inc.
 DOCUMENT TYPE: Journal
 LANGUAGE: English

AB Several brush-type chiral stationary phases (CSPs) based on undecanoyl- or butanoyl-bound (R,R)-1,2-diphenylethane-1,2-diamine (DPEDA) as chiral selector were prepared by an innovative, fast, and less expensive kind of preparation. The key to this method is the immobilization of the enantiomeric pure diamine with only one amino function in a simple substitution reaction on hydroxysuccinimide ester-activated silica. No excess chiral material is lost. Loading can be easily monitored analyzing the filtrate. The free 2nd amino function can subsequently be acylated with different acyl halides. Examples with benzoyl- and 3,5-dinitrobenzoyl (DNB) amides show that, based on the authors' new approach, a library of differently acylated Pirkle-type CSPs can easily be obtained. A benzoylated analog of the com. available ULMO CSP is very effective in separating enantiomers of N-acyl amino acids.

IT 139764-32-2P
 (in-situ preparation of brush-type chiral stationary phases based on undecanoyl- or butanoyl-bound (R,R)-1,2-diphenylethane-1,2-diamine as chiral selector)

RN 139764-32-2 HCAPLUS

CN 2,5-Pyrrolidinedione, 1-[[11-(chlorodimethylsilyl)-1-oxoundecyl]oxy]- (9CI) (CA INDEX NAME)



CC 80-4 (Organic Analytical Chemistry)

IT Silica gel, analysis
 (LiChrospher Si 100; in-situ preparation of brush-type chiral stationary phases based on undecanoyl- or butanoyl-bound (R,R)-1,2-diphenylethane-1,2-diamine as chiral selector)

IT Alcohols, analysis
 Amides, analysis
 (analytes; in-situ preparation of brush-type chiral stationary phases based on undecanoyl- or butanoyl-bound (R,R)-1,2-diphenylethane-1,2-diamine as chiral selector)

IT HPLC stationary phases
 (chiral; in-situ preparation of brush-type chiral stationary phases based on undecanoyl- or butanoyl-bound (R,R)-1,2-diphenylethane-1,2-diamine as chiral selector)

IT Resolution (separation)
 (chromatog.; in-situ preparation of brush-type chiral stationary phases based on undecanoyl- or butanoyl-bound (R,R)-1,2-diphenylethane-1,2-diamine as chiral selector)

IT Amino acids, analysis
 (compds., amide derivative analytes; in-situ preparation of brush-type chiral stationary phases based on undecanoyl- or

butanoyl-bound (R,R)-1,2-diphenylethane-1,2-diamine as chiral selector)

IT Silica gel, analysis
(reaction products; in-situ preparation of brush-type chiral stationary phases based on undecanoyl- or butanoyl-bound (R,R)-1,2-diphenylethane-1,2-diamine as chiral selector)

IT 98-85-1, 1-Phenylethyl alcohol 529-33-9 602-09-5,
1,1'-Bis-2-naphthol 614-14-2, 1-Phenylbutanol
848-75-9 1205-02-3, N-Benzoyl-DL-alanine 1445-91-6,
(S)-1-Phenylethyl alcohol 1466-83-7,
N-Benzoyl-L-leucine 1517-69-7, (R)-1-Phenylethyl alcohol
1517-72-2 1824-74-4 2043-38-1 2198-64-3, N-Benzoyl-L-alanine
2566-22-5, N-Benzoyl-L-phenylalanine 2901-76-0,
N-Benzoyl-DL-phenylalanine 4385-97-1 6296-95-3 7228-47-9
7512-20-1 15914-84-8 17207-57-7 17966-60-8,
N-Benzoyl-D-alanine 17966-67-5, N-Benzoyl-DL-leucine
18531-94-7, (R)-1,1'-Bis-2-naphthol 18531-99-2,
(S)-1,1'-Bis-2-naphthol 22135-49-5 22144-60-1 23357-45-1
26807-65-8 27544-18-9 37002-52-1, N-Benzoyl-D-phenylalanine
42177-25-3 52193-85-8 53732-47-1 57357-55-8,
N-Benzoyl-D-leucine 66719-03-7 67942-91-0 68399-22-4
68399-23-5 69632-33-3 69632-34-4 70622-99-0 74927-93-8
74928-55-5, N-(3,5-DinitroBenzoyl)-DL-phenylalanine 77083-52-4
77083-53-5 83037-88-1, N-(3,5-DinitroBenzoyl)-L-phenylalanine
86091-63-6, N-(3,5-Dinitrobenzoyl)-L-serine 86091-64-7,
N-(3,5-Dinitrobenzoyl)-DL-serine 87068-75-5 90697-07-7
90697-07-7, N-(3,5-Dinitrobenzoyl)-L-isoleucine 94942-50-4
95061-46-4 96782-77-3 96783-07-2 103238-71-7,
N-(3,5-Dinitrobenzoyl)-L-proline 108998-83-0 113679-54-2
113679-56-4 114128-93-7 117466-93-0 119994-29-5
120055-45-0 120055-56-3 120932-64-1, N-(3,5-DinitroBenzoyl)-D-phenylalanine 121758-19-8 127413-52-9 135213-35-3,
N-(3,5-Dinitrobenzoyl)-D-isoleucine 136568-67-7 137036-01-2
143492-62-0, N-(3,5-Dinitrobenzoyl)-DL-proline 143492-63-1,
N-(3,5-Dinitrobenzoyl)-D-proline 143492-64-2,
N-(3,5-Dinitrobenzoyl)-D-serine 143585-47-1 160235-28-9
160333-85-7 160333-86-8 160334-27-0 160334-28-1
170709-41-8 172173-67-0 207460-14-8 207460-15-9
212555-28-7 228114-26-9 228114-27-0 228114-28-1
262844-42-8 263247-50-3 304663-18-1 388091-55-2
388091-57-4 388091-58-5 388099-29-4
(analyte; in-situ preparation of brush-type chiral stationary phases based on undecanoyl- or butanoyl-bound (R,R)-1,2-diphenylethane-1,2-diamine as chiral selector)

IT 999-97-3, 1,1,1,3,3,3-Hexamethyldisilazane
(endcapping reagent; in-situ preparation of brush-type chiral stationary phases based on undecanoyl- or butanoyl-bound (R,R)-1,2-diphenylethane-1,2-diamine as chiral selector)

IT 388091-49-4DP, reaction product with silica gel and end-capped with hexamethyldisilazane 388091-50-7DP, reaction product with silica gel and end-capped with hexamethyldisilazane 388091-51-8DP, reaction products with silica gel and end-capped with hexamethyldisilazane 388091-52-9DP, reaction products with silica gel and end-capped with hexamethyldisilazane
(in-situ preparation of brush-type chiral stationary phases based on undecanoyl- or butanoyl-bound (R,R)-1,2-diphenylethane-1,2-diamine as chiral selector)

IT 98-88-4, Benzoyl chloride 99-33-2, 3,5-Dinitrobenzoyl chloride

112-38-9, 10-Undecenoic acid 625-38-7, 3-Butenoic acid
 1066-35-9, Chlorodimethylsilane 6066-82-6, N-Hydroxysuccinimide
 35132-20-8, (R,R)-1,2-Diphenylethane-1,2-diamine 155075-98-2,
 Kromasil 100 157065-54-8, (R,R)-N-Mono-3,5-dinitrobenzoyl-1,2-
 diphenylethane-1,2-diamine

(in-situ preparation of brush-type chiral stationary
 phases based on undecanoyl- or butanoyl-bound
 (R,R)-1,2-diphenylethane-1,2-diamine as chiral selector)

IT 110661-49-9P, Succinimidyl 10-undecenoate 139764-32-2P
 388091-45-0P, Succinimidyl 3-butenoate 388091-46-1P
 (in-situ preparation of brush-type chiral stationary
 phases based on undecanoyl- or butanoyl-bound
 (R,R)-1,2-diphenylethane-1,2-diamine as chiral selector)
 IT 388610-78-4, ULMO 388610-79-5, ULMO-C 5
 (stationary phase for comparison; in-situ preparation of
 brush-type chiral stationary phases based on undecanoyl- or
 butanoyl-bound (R,R)-1,2-diphenylethane-1,2-diamine as chiral
 selector)

REFERENCE COUNT: 16 THERE ARE 16 CITED REFERENCES AVAILABLE
 FOR THIS RECORD. ALL CITATIONS AVAILABLE
 IN THE RE FORMAT

L15 ANSWER 14 OF 37 HCPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2001:662068 HCPLUS

DOCUMENT NUMBER: 135:359155

TITLE: Ultra-thin coatings of polyvinyl
 alcohol deposited on organic
 monolayers

AUTHOR(S): Kozlov, Mikhail; McCarthy, Thomas J.

CORPORATE SOURCE: Department of Polymer Science and Engineering,
 University of Massachusetts at Amherst,
 Amherst, MA, 01003, USA

SOURCE: Polymer Preprints (American Chemical Society,
 Division of Polymer Chemistry) (2001
), 42(2), 328-329

PUBLISHER: American Chemical Society, Division of Polymer
 Chemistry

DOCUMENT TYPE: Journal; (computer optical disk)

LANGUAGE: English

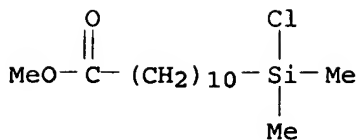
AB Deposition of PVOH on organic monolayers has been found to be
 strongly dependent on the nature of the substrate and the solution
 conditions. PVOH coatings are formed preferably on more
 hydrophobic surfaces, whereas hydrophilic substrates such as
 PEO-terminated monolayers do not support the formation of PVOH
 coatings. Salt content in the solution has been shown to have a
 twofold effect, enhancing hydrophobic interactions and thus
 promoting adsorption, but at the same time impeding aggregation of
 adsorbed polymer by means of hydrogen bonding.

IT 53749-38-5D, 10-(Carbomethoxy)decyldimethylchlorosilane,
 reaction products with silica

(preparation and characterization of ultrathin coatings of
 polyvinyl alc. deposited on organic monolayers)

RN 53749-38-5 HCPLUS

CN Undecanoic acid, 11-(chlorodimethylsilyl)-, methyl ester (9CI)
 (CA INDEX NAME)



CC 42-4 (Coatings, Inks, and Related Products)
 ST ultrathin coating polyvinyl alc deposited org monolayer
 IT Polyoxyalkylenes, uses
 (functionalized, monolayers; preparation and
 characterization of ultrathin coatings of polyvinyl alc
 . deposited on organic monolayers)
 IT Monolayers
 (perfluorinated; preparation and characterization of
 ultrathin coatings of polyvinyl alc. deposited on
 organic monolayers)
 IT Coating materials
 Hydrogen bond
 (preparation and characterization of ultrathin coatings of
 polyvinyl alc. deposited on organic monolayers)
 IT Coupling agents
 (silanes, silica surface modified with; preparation and
 characterization of ultrathin coatings of polyvinyl alc
 . deposited on organic monolayers)
 IT Contact angle
 (water; preparation and characterization of ultrathin
 coatings of polyvinyl alc. deposited on organic
 monolayers)
 IT 7732-18-5, Water, uses
 (contact angle; preparation and characterization of
 ultrathin coatings of polyvinyl alc. deposited on
 organic monolayers)
 IT 7647-14-5, Sodium chloride, uses
 (medium solns.; preparation and characterization of
 ultrathin coatings of polyvinyl alc. deposited on
 organic monolayers)
 IT 25322-68-3D, PEO, functionalized
 (monolayers; preparation and characterization of ultrathin
 coatings of polyvinyl alc. deposited on organic
 monolayers)
 IT 7631-86-9D, Silica, silylated, uses 38051-57-9D,
 n-Decyldimethylchlorosilane, reaction products with
 silica 53749-38-5D, 10-(Carbomethoxy)decyldimethylchloro
 silane, reaction products with silica 102488-47-1D,
 (Tridecafluoro-1,1,2,2-tetrahydrooctyl)dimethylchlorosilane,
 reaction products with silica
 (preparation and characterization of ultrathin coatings of
 polyvinyl alc. deposited on organic monolayers)
 IT 9002-89-5P, Poly(vinyl alcohol)
 (preparation and characterization of ultrathin coatings of
 polyvinyl alc. deposited on organic monolayers)

REFERENCE COUNT: 8 THERE ARE 8 CITED REFERENCES AVAILABLE
 FOR THIS RECORD. ALL CITATIONS AVAILABLE
 IN THE RE FORMAT

L15 ANSWER 15 OF 37 HCPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2000:94047 HCPLUS

DOCUMENT NUMBER: 132:222570

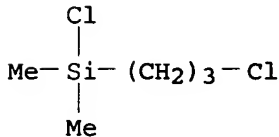
TITLE: The γ -Silicon Effect. IV. The Solvolysis Mechanism of 3-(Aryldimethylsilyl)propyl p-Toluenesulfonates
 AUTHOR(S): Nakashima, Tohru; Fujiyama, Ryoji; Kim, Hyun-Joong; Fujio, Mizue; Tsuno, Yuho
 CORPORATE SOURCE: Inst. Fundamental Res. Org. Chem., Kyushu University, Hakozaki, Fukuoka, 812-8581, Japan
 SOURCE: Bulletin of the Chemical Society of Japan (2000), 73(2), 429-438
 CODEN: BCSJA8; ISSN: 0009-2673
 PUBLISHER: Chemical Society of Japan
 DOCUMENT TYPE: Journal
 LANGUAGE: English

AB Solvolysis rates of 3-(aryldimethylsilyl)propyl p-toluenesulfonates were determined in various solvents. The reaction mechanism of this simple γ -silyl system was clarified based on the solvent effect and the substituent effect analyses. The solvent effect on this system clearly showed the nucleophilic assistance of solvent, but failed to correlate linearly with the extended Winstein-Grunwald equation, substantiating that the reaction should not proceed through either the formation of the cation intermediate or the S_N2 mechanism. This suggests that the reaction takes place in competition between γ -silyl-assisted (k_{Si}) and solvent-assisted (k_s) pathways, and that the competition ratio varies with solvents and with aryl substituents. Product anal. revealed that the former pathway gave only cyclopropane and the latter gave only the substitution products. The overall k_t value could be dissected into the partial rate consts. k_{Si} and k_s for the two pathways by using product ratios. The effects of aryl substituents at the γ -silyl atom on k_{Si} pathway were correlated with unexalted σ^0 parameter, giving the ρ values of -1.0 in 60E (60 volume% ethanol-water) and -1.32 in 97Tw (97 weight% 2,2,2-trifluoroethanol-water), and reflecting the delocalization of incipient carbocationic charge by participation of the Si-C₃ bond. The substituent effects on the k_s pathway were negligibly small; this is in line with the remote reaction center in the concerted S_N2 mechanism.

IT 10605-40-0, 3-(Chlorodimethylsilyl)propyl chloride (metathesis with bromoarene Grignard reagents to give (arylsilyl)propyl chlorides)

RN 10605-40-0 HCAPLUS

CN Silane, chloro(3-chloropropyl)dimethyl- (6CI, 7CI, 8CI, 9CI) (CA INDEX NAME)



CC 29-6 (Organometallic and Organometalloidal Compounds)
ST solvolysis mechanism kinetics arylsilylpropyl tosylate; solvent effect solvolysis arylsilylpropyl tosylate; substituent effect solvolysis arylsilylpropyl tosylate; arylsilylpropyl tosylate prepn solvolysis kinetics mechanism
IT **Alcoholysis**
 Alcoholysis kinetics

Solvolysis**Solvolysis kinetics**

(of (arylsilyl)propyl toluenesulfonates)

IT 75-19-4P, Cyclopropane 261381-52-6P, (3-Ethoxypropyl) (4-methoxyphenyl)dimethylsilane 261381-53-7P, (3-Ethoxypropyl)dimethyl(phenyl)silane 261381-54-8P, (3-Ethoxypropyl)dimethyl(3-(trifluoromethyl)phenyl)silane 261381-55-9P, (3-Ethoxypropyl)dimethyl(4-tolyl)silane 261381-56-0P, (3-Ethoxypropyl)dimethyl(3-tolyl)silane 261381-57-1P, (3-Chloro-4-methoxyphenyl) (3-ethoxypropyl)dimethylsilane 261381-58-2P, (4-Chlorophenyl) (3-ethoxypropyl)dimethylsilane 261381-59-3P, Dimethyl(3-(2,2,2-trifluoroethoxy)propyl) (3-(trifluoromethyl)phenyl)silane (formation in **ethanolysis of (arylsilyl)propyl tosylate**)

IT 10605-40-0, 3-(Chlorodimethylsilyl)propyl chloride (metathesis with bromoarene Grignard reagents to give (arylsilyl)propyl chlorides)

IT 98-59-9, p-Toluenesulfonyl chloride (metathesis with lithiated (arylsilyl)propanols)

IT 57292-94-1P, 3-(Dimethyl(4-tolyl)silyl)-1-propanol 68469-62-5P, 3-(Dimethyl(phenyl)silyl)-1-propanol 68469-63-6P, 3-((4-Methoxyphenyl)dimethylsilyl)-1-propanol 261381-42-4P, 3-((3-Chloro-4-methoxyphenyl)dimethylsilyl)-1-propanol 261381-43-5P, 3-(Dimethyl(3-tolyl)silyl)-1-propanol 261381-44-6P, 3-((4-Chlorophenyl)dimethylsilyl)-1-propanol 261381-45-7P, 3-(Dimethyl(3-(trifluoromethyl)phenyl)silyl)-1-propanol 261381-46-8P, 1,1-Dideutero-3-(dimethyl(phenyl)silyl)-1-propanol 261381-47-9P, 1,1-Dideutero-3-((4-methoxyphenyl)dimethylsilyl)-1-propanol 261381-48-0P, 1,1-Dideutero-3-(dimethyl(3-(trifluoromethyl)phenyl)silyl)-1-propanol (preparation and condensation with tosyl chloride)

IT 2632-95-3P, (3-Chloropropyl)dimethyl(phenyl)silane 54040-86-7P, (3-Chloropropyl) (4-methoxyphenyl)dimethylsilane 54040-87-8P, (3-Chloropropyl)dimethyl(4-tolyl)silane 123015-76-9P, (3-Chloropropyl)dimethyl(3-(trifluoromethyl)phenyl)silane 123015-78-1P, (4-Chlorophenyl) (3-chloropropyl)dimethylsilane 130284-15-0P, (3-Chloropropyl)dimethyl(3-tolyl)silane 261381-41-3P, (3-Chloro-4-methoxyphenyl) (3-chloropropyl)dimethylsilane (preparation and conversion to alc. via Grignard reagent and oxygen)

IT 167282-62-4P, 3-(Dimethyl(phenyl)silyl)propyl 4-methylbenzenesulfonate 261381-35-5P, 3-((4-Methoxyphenyl)dimethylsilyl)propyl 4-methylbenzenesulfonate 261381-36-6P, 3-((3-Chloro-4-methoxyphenyl)dimethylsilyl)propyl 4-methylbenzenesulfonate 261381-37-7P, 3-(Dimethyl(4-tolyl)silyl)propyl 4-methylbenzenesulfonate 261381-38-8P, 3-(Dimethyl(3-tolyl)silyl)propyl 4-methylbenzenesulfonate 261381-39-9P, 3-((4-Chlorophenyl)dimethylsilyl)propyl 4-methylbenzenesulfonate 261381-40-2P, 3-(Dimethyl(3-(trifluoromethyl)phenyl)silyl)propyl 4-methylbenzenesulfonate 261381-49-1P, 1,1-Dideutero-3-(dimethyl(phenyl)silyl)propyl 4-methylbenzenesulfonate 261381-50-4P, 1,1-Dideutero-3-((4-methoxyphenyl)dimethylsilyl)propyl 4-methylbenzenesulfonate 261381-51-5P, 1,1-Dideutero-3-(dimethyl(3-(trifluoromethyl)phenyl)silyl)propyl 4-methylbenzenesulfonate (preparation and kinetics and mechanism of solvolysis of)

REFERENCE COUNT: 42 THERE ARE 42 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

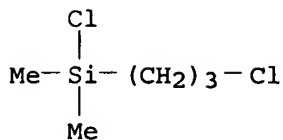
L15 ANSWER 16 OF 37 HCPLUS COPYRIGHT 2006 ACS on STN
 ACCESSION NUMBER: 1999:680913 HCPLUS
 DOCUMENT NUMBER: 132:56942
 TITLE: Communication between Surfaces by Electron Relay in a Doubly Heterogeneous Photochemical Reaction
 AUTHOR(S): Ayadim, Mohamed; Jiwan, Jean L. Habib; Soumillion, Jean Ph.
 CORPORATE SOURCE: Laboratory of Photochemistry, Catholic University of Louvain, Louvain La Neuve, B-1348, Belg.
 SOURCE: Journal of the American Chemical Society (1999), 121(44), 10436-10437
 CODEN: JACSAT; ISSN: 0002-7863
 PUBLISHER: American Chemical Society
 DOCUMENT TYPE: Journal
 LANGUAGE: English

AB It is shown that the surface of organically modified silica beads may be photoactivated using a sensitizer in the solution together with an electron relay: an amine is unhooked from a sulfonamide substrate attached to the silica. Furthermore it was found that a doubly heterogeneous sensitized system in which the sensitizer and the sulfonamide substrate are attached to different silica beads works even better: an electron relaying shuttle ensures communication between the beads. This surface to surface communication is, to the authors knowledge, the first example of a photoinduced electron transfer organic reaction photosensitized on one surface and relayed to another one. A nonphotochem. system related to the authors work may be cited: an alc. was shown to be alternately reduced and oxidized on the surfaces of silica and alumina beads charged with appropriate redox reagents (Kim, B. et al., 1983). In this paper the photoreductive deprotection of sulfonamides is taken as a test reaction for the study of heterogeneous conditions: in the authors example, 1,4-dimethoxynaphthalene sensitizes the cleavage of the sulfonamides in the presence of potassium borohydride coreductant. Four different exptl. systems were used: homogeneous reaction (system A), heterogeneous system with sensitizer (system B) or substrate (system C) covalently grafted on silica, and the doubly heterogeneous system with sensitizer and substrate grafted on different silica beads (system D).

IT 10605-40-0, 3-Chloropropyldimethylchlorosilane (photoreactions in heterogeneous systems containing photosensitizer and sulfonamide reactants grafted on different silica beads and electron relaying shuttle ensuring communication between these beads)

RN 10605-40-0 HCPLUS

CN Silane, chloro(3-chloropropyl)dimethyl- (6CI, 7CI, 8CI, 9CI) (CA INDEX NAME)



CC 74-1 (Radiation Chemistry, Photochemistry, and Photographic and Other Reprographic Processes)

IT 91-20-3, Naphthalene, processes
(electron relay; photoreactions in heterogeneous systems containing photosensitizer and sulfonamide reactants grafted on different silica beads and electron relaying shuttle ensuring communication between these beads)

IT 10605-40-0, 3-Chloropropyldimethylchlorosilane
(photoreactions in heterogeneous systems containing photosensitizer and sulfonamide reactants grafted on different silica beads and electron relaying shuttle ensuring communication between these beads)

IT 7631-86-9D, Silica, surface modified, processes
(photoreactions in heterogeneous systems containing photosensitizer and sulfonamide reactants grafted on different silica beads and electron relaying shuttle ensuring communication between these beads)

IT 5450-75-9D, surface reaction product with chloropropyl-modified silica or trimethylated silica
(photoreactions in heterogeneous systems containing photosensitizer and sulfonamide reactants grafted on different silica beads and electron relaying shuttle ensuring communication between these beads)

IT 252848-35-4DP, surface reaction product with chloropropyl-modified or trimethylated silica
(photoreactions in heterogeneous systems containing photosensitizer and sulfonamide reactants grafted on different silica beads and electron relaying shuttle ensuring communication between these beads)

IT 84-85-5D, 4-Methoxy-1-naphthol, surface reaction product with chloropropyl-modified or trimethylsilylated silica
(photosensitizer; photoreactions in heterogeneous systems containing photosensitizer and sulfonamide reactants grafted on different silica beads and electron relaying shuttle ensuring communication between these beads)

IT 75-77-4, Trimethylchlorosilane, uses 121-44-8, Triethylamine, uses
(preparation of trimethylsilylated silica surface)

IT 114958-23-5P
(reaction with dicyclohexylcarbodiimide and hydroxysuccinimide in synthesis of sulfonamide derivative)

REFERENCE COUNT: 14 THERE ARE 14 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

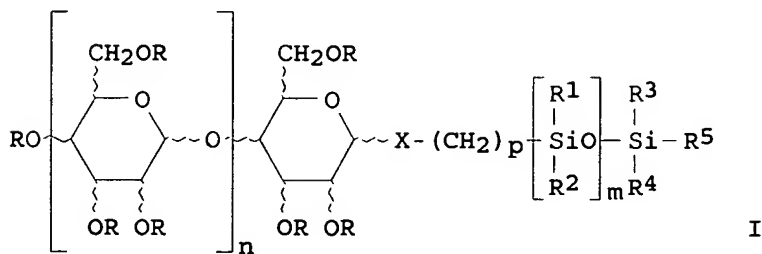
L15 ANSWER 17 OF 37 HCPLUS COPYRIGHT 2006 ACS on STN
 ACCESSION NUMBER: 1999:228013 HCPLUS
 DOCUMENT NUMBER: 130:312019
 TITLE: Preparation of organopolysiloxane compounds having sugar residues as dermal absorption enhancers for drugs
 INVENTOR(S): Nagase, Hiroshi; Akimoto, Satoko; Aoyagi,

PATENT ASSIGNEE(S): Takao; Akiyama, Eiichi
 SOURCE: Sagami Chemical Research Center, Japan
 Jpn. Kokai Tokkyo Koho, 17 pp.
 CODEN: JKXXAF

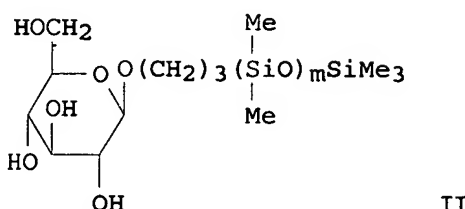
DOCUMENT TYPE: Patent
 LANGUAGE: Japanese
 FAMILY ACC. NUM. COUNT: 1
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 11092490	A2	19990406	JP 1998-200151	1998 0715
PRIORITY APPLN. INFO.:			JP 1997-200322	A 1997 0725
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GI



I



II

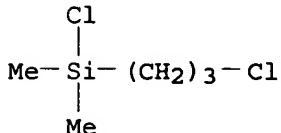
AB The polysiloxane glycosides (I; R = H, acyl; X = O, S; R1-R4 = C1-6 alkyl; R5 = C1-20 alkyl; n = 0,1,2; p = 3-6; m = integer of ≥ 1) are prepared. Also claimed is a dermal absorption enhancer containing I (R = H; X, R1-R5, n,p,m = same as above) for drugs. The above compds. I possess good dermal absorption-enhancing effect not only for hydrophobic but also water-soluble drugs and are low in skin-irritation and toxicity and are used in a drug delivery system. Thus, allyl 2,3,4,6-tetraacetyl- β -D-glucopyranoside (preparation given) underwent add. reaction (hydrosilylation) with H(SiMe2O)mSiMe3 in the presence of dicyclopentadienyl platinum dichloride in THF at 70° for 2 h followed by deacetylation

with NaOMe in MeOH gave the title compound (II; m = integer of ≥ 1). A solution containing 20 mg antipyrin (antiinflammatory agent) and 2 weight% II in 2 mL 50% aqueous EtOH in a donor chamber was contacted through a rabbit abdominal skin with a solution of a phosphate buffer (pH 7.4) in a receptor chamber in a 2-chamber cell at 37° for 12 h while both chambers were stirred. The cumulative amount of antipyrin permeated through the skin was 0.189 and 0.935 mg after 6 and 12 h, resp., vs. 0.056 and 0.140 mg after 6 and 12 h, resp.

IT 10605-40-0, (3-Chloropropyl)dimethylsilyl chloride
(preparation of organopolysiloxane compds. having sugar residues as dermal absorption enhancers for drugs)

RN 10605-40-0 HCAPLUS

CN Silane, chloro(3-chloropropyl)dimethyl- (6CI, 7CI, 8CI, 9CI) (CA INDEX NAME)



IC ICM C07H015-04
ICS C07H015-04; A61K047-26; A61K047-30

CC 33-4 (Carbohydrates)
Section cross-reference(s): 1

ST polysiloxane glycoside prep drug delivery system;
organopolysiloxane contg sugar prep drug dermal absorption enhancer

IT Drug delivery systems
(carriers; preparation of organopolysiloxane compds.
having sugar residues as dermal absorption enhancers for drugs)

IT Drug delivery systems
(preparation of organopolysiloxane compds. having sugar residues as dermal absorption enhancers for drugs)

IT Polysiloxanes, preparation
(preparation of organopolysiloxane compds. having sugar residues as dermal absorption enhancers for drugs)

IT 157622-01-0P 223536-19-4P 223536-21-8P 223536-23-0P
(preparation of organopolysiloxane compds. having sugar residues as dermal absorption enhancers for drugs)

IT 62-56-6, Thiourea, reactions 107-18-6, 2-Propen-1-ol, reactions 112-41-4, 1-Dodecene 604-69-3, β -D-Glucose pentaacetate 1066-35-9, Dimethylsilyl chloride 1066-40-6, Trimethylsilanol 3277-26-7 6919-96-6, 2,3,4,6-Tetra-O-acetyl- β -D-glucopyranosyl bromide 10605-40-0, (3-Chloropropyl)dimethylsilyl chloride 20764-63-0, D-(+)-Cellobiose octaacetate
(preparation of organopolysiloxane compds. having sugar residues as dermal absorption enhancers for drugs)

IT 10343-15-4P, Allyl 2,3,4,6-tetra-O-acetyl- β -D-glucopyranoside 40591-65-9P, S-(2,3,4,6-Tetra-O-acetyl- β -D-glucopyranosyl)isothiourea hydrobromide 50256-34-3P 128147-45-5P 172413-82-0P 172413-83-1P 223536-24-1P 223536-25-2P 223536-26-3P 223536-27-4P
(preparation of organopolysiloxane compds. having sugar residues as dermal absorption enhancers for drugs)

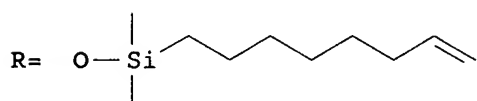
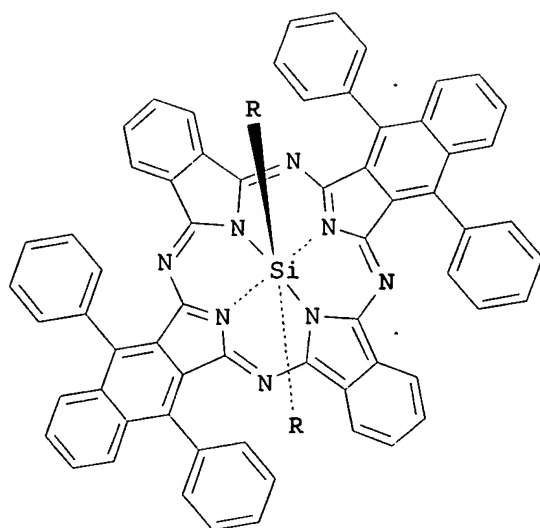
L15 ANSWER 18 OF 37 HCAPLUS COPYRIGHT 2006 ACS on STN
 ACCESSION NUMBER: 1998:392268 HCAPLUS
 DOCUMENT NUMBER: 129:78836
 TITLE: Fluorescence energy transfer and
 intramolecular energy transfer in particles
 using novel compounds for the application in
 immunoassays and nucleic acid assays
 INVENTOR(S): Buechler, Kenneth F.; Noar, J. Barry; Tadesse,
 Lema
 PATENT ASSIGNEE(S): Biosite Diagnostics Inc., USA
 SOURCE: U.S., 36 pp., Cont.-in-part of U. S. Ser. No.
 274,534.
 CODEN: USXXAM
 DOCUMENT TYPE: Patent
 LANGUAGE: English
 FAMILY ACC. NUM. COUNT: 7
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 5763189	A	19980609	US 1994-311098	1994 0923
US 6238931	B1	20010529	US 1994-274534	1994 0712
US 6251687	B1	20010626	US 1995-409298	1995 0323
US 5824799	A	19981020	US 1996-620597	1996 0322
US 6964844	B1	20051115	US 1998-66255	1998 0424
US 2002061602	A1	20020523	US 2001-776599	2001 0201
PRIORITY APPLN. INFO.:			US 1993-126367	B2 1993 0924
			US 1993-138708	B2 1993 1018
			US 1994-274534	A2 1994 0712
			US 1994-311098	A2 1994 0923

<--	WO 1994-US10826	W	1994
			0923
<--	US 1995-409298	A2	1995
			0323
<--	US 1995-409825	A2	1995
			0323
<--	US 1996-620597	A1	1996
			0322
<--	US 1998-66255	A2	1998
			0424
<--			

OTHER SOURCE(S) :
GI

MARPAT 129:78836



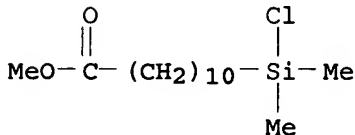
AB The invention concerns the **synthesis** of novel dyes and methods for the detection or visualization of analytes; more specifically fluorescent latex particles which randomly incorporate the novel fluorescent dyes and utilize fluorescent energy transfer and intramol. energy transfer, for the detection

of analytes in immunoassays or in nucleic acid assay. Particles comprise an energy donor as a first component and a fluorescent dye as a second component that are positioned at an energy exchanging distance from one another; the two components have a Stokes shift of greater than or equal to 50 nm; and the particles bind on the surface a protein, polypeptide, nucleic acid, nucleotide or protein containing ligand analog. In addition, novel fluorescent dyes (e.g., I) are described which exhibit intramol. energy transfer for use to label various mols., proteins, polypeptides, nucleotides and nucleic acids or to incorporate into particles. Compns. are given to minimize fluorescence quenching and to maximize fluorescence intensities of the dye mols. in the particles through the use of different dye mols. which posses the same or very similar excitation and emission wavelengths. Many novel phthalocyanine derivs. and hybrid phthalocyanine derivs. are disclosed. Thus latex microparticles have at least one hybrid phthalocyanine derivative, that derivative has at least one donor subunit with a desired excitation peak; and at least one acceptor unit with desired emission peak. The derivative(s) is/are capable of intramol. energy transfer from the donor subunit to the acceptor subunit; such derivs. also may contain and electron transfer subunit. Axial ligands may covalently bound to the metals contained in the hybrid phthalocyanine derivs. Numerous compds. capable of intramol. energy transfer as well as compds. for fluorescence energy transfer were synthesized.

IT 53749-38-5, (10-Carbomethoxydecyl)dimethylchlorosilane
(Fluorescence energy transfer and intramol. energy transfer in particles using novel compds. for the application in immunoassays and nucleic acid assays)

RN 53749-38-5 HCAPLUS

CN Undecanoic acid, 11-(chlorodimethylsilyl)-, methyl ester (9CI)
(CA INDEX NAME)



IC ICM G01N033-542
ICS G01N033-543; C09K009-00; C09K009-02

INCL 435007100

CC 9-16 (Biochemical Methods)
Section cross-reference(s): 3, 15, 28, 29

ST fluorescence dye energy transfer latex immunoassay; phthalocyanine naphthalocyanine deriv **synthesis** fluorescent dye

IT 68-12-2P, Dimethylformamide, **preparation** 71-41-0P, 1-Pentanol, **preparation** 109-99-9P, Tetrahydrofuran, **preparation** 119-64-2P, 1,2,3,4-Tetrahydronaphthalene
(Fluorescence energy transfer and intramol. energy transfer in particles using novel compds. for the application in immunoassays and nucleic acid assays)

IT 75-78-5, Dichlorodimethylsilane 76-86-8, Triphenylchlorosilane 597-52-4, Triethylsilanol 1631-83-0, Diphenylchlorosilane 1719-58-0, Chlorodimethylvinylsilane 1835-65-0, Tetrafluorophthalonitrile 3468-11-9, 1,3-Diiminoisoindoline 3634-67-1, Chlorotrihexylsilane

6554-98-9, trans-4-Hydroxystilbene 7646-78-8, Tin tetrachloride, reactions 10026-04-7, Silicon tetrachloride 10038-98-9, Germanium tetrachloride 10264-67-2 17196-12-2, 7-Oct-enyldimethylchlorosilane 18156-15-5, Chloro(3-cyanopropyl)-dimethylsilane 19333-10-9, Silicon phthalocyanine dichloride 20082-71-7, Chlorodimethylpentafluorophenylsilane 26857-61-4 32703-80-3, 4-tert-Butylphthalonitrile 37623-03-3, 1,4-Diphenylnaphthalene-2,3-dicarbonitrile 53749-38-5, (10-Carbomethoxydecyl)dimethylchlorosilane 74815-81-9, 2,3-Dibromo-6,7-dicyanonaphthalene 92396-91-3 102488-47-1 105528-25-4 116453-89-5, 1,4-Dibutoxynaphthalene-2,3-dicarbonitrile 183872-68-6, 4,7-Diethoxy-1,3-diiminoisoindoline (Fluorescence energy transfer and intramol. energy transfer in particles using novel compds. for the application in immunoassays and nucleic acid assays)

REFERENCE COUNT: 21 THERE ARE 21 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L15 ANSWER 19 OF 37 HCPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1998:65914 HCPLUS

DOCUMENT NUMBER: 128:115084

TITLE: Functionalized ferrocenyldiphosphines, a process for their preparation and their use

INVENTOR(S): Pugin, Benoit; Landert, Heidi

PATENT ASSIGNEE(S): Novartis A.-G., Switz.; Pugin, Benoit; Landert, Heidi

SOURCE: PCT Int. Appl., 91 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 9801457	A1	19980115	WO 1997-EP3626	1997 0709

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W: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GE, GH, HU, IL, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM

RW: GH, KE, LS, MW, SD, SZ, UG, ZW, AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG

CA 2256770	AA	19980115	CA 1997-2256770	1997 0709
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AU 9736211	A1	19980202	AU 1997-36211	1997 0709
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EP 912586	A1	19990506	EP 1997-932789
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EP 912586 B1 20020116
 R: AT, BE, CH, DE, ES, FR, GB, IT, LI, NL
 JP 2000514436 T2 20001031 JP 1998-504804

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 AT 212032 E 20020215 AT 1997-932789

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 ES 2171267 T3 20020901 ES 1997-932789

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 US 6169192 B1 20010102 US 1999-214667

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 PRIORITY APPLN. INFO.: CH 1996-1746

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 CH 1996-2069

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 WO 1997-EP3626

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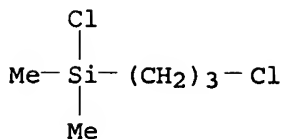
OTHER SOURCE(S): MARPAT 128:115084

AB The invention relates to 1,2-ferrocenyldiphosphines which contain a functional group in the 1' position either directly or via a bridging group, and also a process for their preparation. The compds. are important ligands for transition metal complexes containing d-8 metals such as Rh, Ru, Pd or Ir. These transition metal complexes are widely used in the hydrogenation of organic double or triple bonds, in particular olefinic double bonds and C-heteroatom double bonds. The complexes are particularly suitable for enantioselective hydrogenation using chiral ferrocenyldiphosphines and corresponding prochiral unsatd. compds. Ferrocenyldiphosphines having a functional group in the 1' position are also important intermediates for ferrocenyldiphosphine ligands and their metal complexes of d-8 metals such as Rh, Ru, Pd or Ir which are bound to inorg. or organic polymeric supports via this functional group. These metal complexes bound to an inorg. or organic support material are likewise very suitable for the hydrogenation of organic double or triple bonds.

IT 10605-40-0, Chloro(3-chloropropyl)dimethylsilane
 (preparation of ferrocenyldiphosphines as hydrogenation catalysts)

RN 10605-40-0 HCAPLUS

CN Silane, chloro(3-chloropropyl)dimethyl- (6CI, 7CI, 8CI, 9CI) (CA INDEX NAME)



IC ICM C07F017-02
 ICS B01J031-28; C07B031-00; C07B053-00; C08F112-08
 CC 29-12 (Organometallic and Organometalloidal Compounds)
 Section cross-reference(s): 34
 ST ferrocenyl phosphine prepn hydrogenation catalyst
 IT Polyoxyalkylenes, reactions
 (preparation of ferrocenyldiphosphines as hydrogenation catalysts)
 IT Silica gel, reactions
 (reaction products; polymers as supports for ferrocenyldiphosphines hydrogenation catalysts)
 IT 9002-89-5, Polyvinyl alcohol 9003-01-4 9003-70-7D, aminomethylated 25322-68-3
 (preparation of ferrocenyldiphosphines as hydrogenation catalysts)
 IT 201551-71-5P 201552-06-9P
 (preparation of ferrocenyldiphosphines as hydrogenation catalysts)
 IT 201551-65-7P 201551-77-1P 201551-90-8P 201552-03-6DP, polyethyleneglycol (methyl)diisocyanophenylated, supported 201552-06-9DP, Amberlite IRC 50 supported 201552-06-9DP, Amberlite IRC 76 supported 201552-06-9DP, polyacrylic acid supported 201552-11-6DP, (methyl)diisocyanophenylated aminomethylated polystyrene divinylbenzene supported 201552-11-6DP, aminomethylated polystyrene divinylbenzene supported 201552-11-6DP, polyvinyl alc. supported
 (preparation of ferrocenyldiphosphines as hydrogenation catalysts)
 IT 78-95-5, Chloroacetone 85-41-6, Phthalimide 90-80-2
 105-53-3, Diethyl malonate 109-73-9, Butylamine, reactions
 111-36-4, Butyl isocyanate 503-30-0, Oxetane 530-62-1,
 1,1'-Carbonyldimidazole 584-84-9 996-82-7, Diethyl sodiomalonate 1079-66-9, Chlorodiphenylphosphine 4009-98-7, Methoxymethyltriphenylphosphonium chloride 9002-29-3, Amberlite IRC 50 10605-40-0, Chlоро(3-chloropropyl)dimethylsilane 24801-88-5 31886-58-5 71360-06-0, Bis(3,5-xylyl)phosphine 153550-33-5, Amberlite IRC 76 183720-68-5 183720-73-2
 (preparation of ferrocenyldiphosphines as hydrogenation catalysts)
 IT 182227-18-5P 182227-20-9P 201551-64-6P 201551-66-8P
 201551-69-1P 201551-75-9P 201551-88-4P 201551-99-7P
 201552-03-6P 201552-05-8P 201552-08-1P 201552-11-6P
 (preparation of ferrocenyldiphosphines as hydrogenation catalysts)
 IT 201551-68-0P 201551-73-7P 201551-81-7P 201551-92-0P
 201551-94-2P 201551-96-4P 201551-99-7DP, polystyrene supported 201552-01-4P 201552-03-6DP, hydrogenated hydroxy terminated polybutadiene (methyl)diisocyanophenylated, supported 201552-07-0P 201552-09-2P 201552-10-5P
 (preparation of ferrocenyldiphosphines as hydrogenation catalysts)

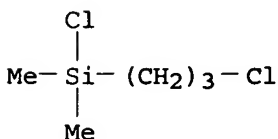
REFERENCE COUNT: 5 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE

IN THE RE FORMAT

L15 ANSWER 20 OF 37 HCPLUS COPYRIGHT 2006 ACS on STN
 ACCESSION NUMBER: 1997:748123 HCPLUS
 DOCUMENT NUMBER: 128:145262
 TITLE: Polymeric percutaneous drug penetration enhancer. *Synthesis* and enhancing property of PEG/PDMS block copolymer with a cationic end group
 AUTHOR(S): Akimoto, Tomoko; Aoyagi, Takao; Minoshima, Jun-ichi; Nagase, Yu
 CORPORATE SOURCE: Sagamihara, Nishi-Ohnuma, Sagami Chemical Research Center, Kanagawa 229, 4-4-1, Japan
 SOURCE: Journal of Controlled Release (1997), 49(2,3), 229-242
 CODEN: JCREEC; ISSN: 0168-3659
 PUBLISHER: Elsevier Science B.V.
 DOCUMENT TYPE: Journal
 LANGUAGE: English

AB Polyethylene glycol/polydimethylsiloxane (PEG/PDMS) block copolymers containing an ammonium moiety at one chain end with various mol. wts. were prepared to develop a silicone-based polymeric transdermal penetration enhancer. As the precursor of the desired block copolymer, 3-chloropropyl-terminated PEG/PDMS block copolymers were prepared via an initiator method, i.e. the anionic ring-opening polymerization of hexamethylcyclotrisiloxane was carried out by initiating with silanolate anion derived from PEG-silanol, α -3-(dimethylhydroxysilyl)propyl- ω -methyl-PEG oligomer. The initiator, PEG-silanol, was obtained from α -allyl-PEG by hydrosilylation with dimethylethoxysilane, followed by hydrolysis of the ethoxysilyl group. The enhancing activity in the drug penetration was evaluated by in vitro expts. using a two-chamber diffusion cell. Indomethacin and antipyrine were used as hydrophobic and hydrophilic model drugs, resp., and the amts. of drugs permeating through the rabbit abdominal skin were measured with or without these polymeric enhancers. These enhancers were very effective for the penetration of hydrophilic drug, but not for that of hydrophobic one. On the other hand, the enhancing activities were influenced by the chain length of PDMS and PEG components. A suitable balance between the hydrophobic PDMS segment and the hydrophilic PEG segment would exist for a high enhancing activity of drug penetration. It was also found that the enhancing activity was due to an increase of the partition coefficient of a drug into the stratum corneum, from the determination of kinetic parameters in the drug permeation.

IT 10605-40-0, 3-Chloropropyldimethylchlorosilane
 (preparation and percutaneous enhancing property of PEG/PDMS block copolymer with a cationic end group)
 RN 10605-40-0 HCPLUS
 CN Silane, chloro(3-chloropropyl)dimethyl- (6CI, 7CI, 8CI, 9CI) (CA INDEX NAME)



CC 63-5 (Pharmaceuticals)
 Section cross-reference(s): 35
 IT Biological transport
 (permeation; preparation and percutaneous enhancing
 property of PEG/PDMS block copolymer with a cationic end group)
 IT Polysiloxanes, biological studies
 (polyoxyethylene-, block, quaternary ammonium group-terminated;
 preparation and percutaneous enhancing property of PEG/PDMS
 block copolymer with a cationic end group)
 IT Skin
 (preparation and percutaneous enhancing property of
 PEG/PDMS block copolymer with a cationic end group)
 IT Drug delivery systems
 (transdermal; preparation and percutaneous enhancing
 property of PEG/PDMS block copolymer with a cationic end group)
 IT 170210-26-1DP, quaternary ammonium-group terminated
 (preparation and percutaneous enhancing property of
 PEG/PDMS block copolymer with a cationic end group)
 IT 598-56-1DP, N,N-Dimethylethylamine, reaction products
 with iodopropyl-terminated PEG-polydimethylsiloxane copolymer
 927-62-8DP, 1-Butanamine, N,N-dimethyl-, reaction products
 with iodopropyl-terminated PEG-polydimethylsiloxane copolymer
 7378-99-6DP, 1-Octanamine, N,N-dimethyl-, reaction
 products with iodopropyl-terminated PEG-
 polydimethylsiloxane copolymer
 (preparation and percutaneous enhancing property of
 PEG/PDMS block copolymer with a cationic end group)
 IT 53-86-1, Indomethacin 60-80-0, Antipyrine
 (preparation and percutaneous enhancing property of
 PEG/PDMS block copolymer with a cationic end group)
 IT 106-95-6, 3-Bromopropene, reactions 9004-74-4 10605-40-0
 , 3-Chloropropyldimethylchlorosilane 14857-34-2,
 Dimethylethoxysilane
 (preparation and percutaneous enhancing property of
 PEG/PDMS block copolymer with a cationic end group)
 IT 27252-80-8P, Poly(oxy-1,2-ethanediyl), α -methyl- ω -(2-
 propenoxy)- 164149-53-5P 202209-58-3P 202209-59-4P
 (preparation and percutaneous enhancing property of
 PEG/PDMS block copolymer with a cationic end group)

REFERENCE COUNT: 17 THERE ARE 17 CITED REFERENCES AVAILABLE
 FOR THIS RECORD. ALL CITATIONS AVAILABLE
 IN THE RE FORMAT

L15 ANSWER 21 OF 37 HCAPLUS COPYRIGHT 2006 ACS on STN
 ACCESSION NUMBER: 1996:761698 HCAPLUS
 DOCUMENT NUMBER: 126:33023
 TITLE: Hybrid phthalocyanine derivatives and their
 uses
 INVENTOR(S): Buechler, Kenneth F.; Noar, Joseph B.;
 Tadesse, Lema
 PATENT ASSIGNEE(S): Biosite Diagnostics Incorporated, USA
 SOURCE: PCT Int. Appl., 190 pp.
 CODEN: PIXXD2
 DOCUMENT TYPE: Patent
 LANGUAGE: English
 FAMILY ACC. NUM. COUNT: 7
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
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WO 9629367 A1 19960926 WO 1996-US3833
1996
0322

W: AL, AM, AT, AU, AZ, BB, BG, BR, BY, CA, CH, CN, CZ, DE,
DK, EE, ES, FI, GB, GE, HU, IS, JP, KE, KG, KP, KR, KZ,
LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ,
PL, PT, RO, RU, SD, SE, SG, SI
RW: KE, LS, MW, SD, SZ, UG, AT, BE, CH, DE, DK, ES, FI, FR,
GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI,
CM, GA, GN, ML

CA 2215727 AA 19960926 CA 1996-2215727
1996
0322

CA 2215727 C 20031230
AU 9653188 A1 19961008 AU 1996-53188
1996
0322

EP 820489 A1 19980128 EP 1996-909805
1996
0322

EP 820489 B1 20010711
R: AT, CH, DE, ES, FR, GB, IT, LI, NL
JP 10508897 T2 19980902 JP 1996-528604
1996
0322

JP 3388753 B2 20030324
AT 203045 E 20010715 AT 1996-909805
1996
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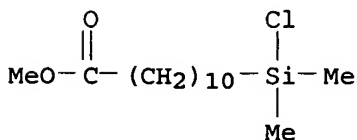
PRIORITY APPLN. INFO.: US 1995-409825 A
1995
0323

WO 1996-US3833 W
1996
0322

AB Water-soluble hybrid phthalocyanine derivs., fluorescent latex particles incorporating which are useful in competitive and noncompetitive immunoassays and nucleic acid assays, have (1) ≥ 1 donor subunit with a desired excitation peak and (2) ≥ 1 acceptor subunit with a desired emission peak, and are capable of intramol. energy transfer from the donor subunit to the acceptor subunit. They may also contain an electron-transfer subunit. Axial ligands may be covalently bound to the metals contained in the water-soluble hybrid phthalocyanine derivs. Ligands, ligand analogs, polypeptides, proteins, and nucleic acids can be linked to the axial ligands of the dyes to form conjugates useful in immunoassays and nucleic acid assays.

IT 53749-38-5, (10-Carbomethoxydecyl)chlorodimethylsilane (preparation of water-soluble fluorescent hybrid phthalocyanine derivs. for immunoassays)

RN 53749-38-5 HCAPLUS
 CN Undecanoic acid, 11-(chlorodimethylsilyl)-, methyl ester (9CI)
 (CA INDEX NAME)



IC ICM C09B047-04
 ICS G01N033-533; C07D487-22; C07F007-08
 ICI C07D487-22, C07D259-00, C07D209-00
 CC 41-7 (Dyes, Organic Pigments, Fluorescent Brighteners, and
 Photographic Sensitizers)
 Section cross-reference(s): 9, 15
 IT Latex
 (anti-human chorionic gonadotropin antibody conjugates;
 preparation of water-soluble fluorescent spitzehybrid
 phthalocyanine derivs. as dye systems for immunoassays)
 IT Immunoassay
 (preparation of water-soluble fluorescent hybrid
 phthalocyanine derivs. as dye systems for immunoassays)
 IT Polyoxyalkylenes, reactions
 (preparation of water-soluble fluorescent hybrid
 phthalocyanine derivs. for immunoassays)
 IT Blood analysis
 Body fluid
 Fluorescent dyes
 (preparation of water-soluble fluorescent spitzehybrid
 phthalocyanine derivs. as dye systems for immunoassays)
 IT Nucleic acids
 (preparation of water-soluble fluorescent spitzehybrid
 phthalocyanine derivs. as dye systems for immunoassays)
 IT Antibodies
 (sulfonated hybrid phthalocyanine derivative conjugates;
 preparation of water-soluble fluorescent hybrid phthalocyanine
 derivs. as dye systems for immunoassays)
 IT 9002-61-3, Chorionic gonadotropin
 (antibodies specific for; preparation of water-soluble
 fluorescent hybrid phthalocyanine derivs. and antibody
 conjugates for immunoassays)
 IT 9003-53-6D, Polystyrene, sulfate
 (latex; preparation of water-soluble fluorescent spitzehybrid
 phthalocyanine derivs. as dye systems for immunoassays)
 IT 67881-06-5P 68812-20-4P 92396-89-9P,
 Bis[(trihexylsilyl)oxy]silicon phthalocyanine 117753-12-5P
 142700-81-0DP, sulfonated 149971-18-6P 153454-01-4P
 163968-88-5P 163968-89-6P 163968-91-0P, Silicon, bis(
 ethenylidiphenylsilanolato) [37H,39H-tetranaphtho[2,3-
 b:2',3'-g:2'',3''-l:2''',3'''-q]porphyrazinato(2-)-
 N37,N38,N39,N40]-, (OC-6-12)- 163968-92-1P 163968-94-3P
 163968-95-4P 163969-00-4P 163969-01-5P 163969-07-1P
 163969-08-2P 163969-09-3P 163969-10-6P 163969-11-7P
 163969-14-0P 163969-15-1P 163969-26-4P 171118-91-5P
 171118-94-8P 183872-48-2P 183872-49-3P 183872-50-6P
 183872-51-7P 183872-55-1P 183872-56-2P 183872-57-3DP,
 sulfonated 183872-57-3P 183872-58-4P 183872-59-5P

183872-61-9P	183872-62-0P	183872-63-1P	183872-66-4P
183872-71-1P	183872-72-2P	183872-74-4P	183872-76-6P
183872-77-7P	183872-79-9P	183872-81-3P	183872-82-4P
183872-84-6P	183872-85-7P	183872-86-8P	183872-87-9P
183872-88-0P	183872-89-1P	183872-90-4P	183872-92-6P
183872-94-8P	183872-95-9P	183872-96-0P	183872-97-1P
183872-98-2P	183872-99-3P	183873-00-9P	183873-03-2P
183873-05-4DP, sulfonated	183873-07-6DP, sulfonated		
183873-11-2DP, sulfonated	183873-13-4DP, sulfonated		
183873-14-5DP, sulfonated	183873-15-6DP, sulfonated		
183873-17-8DP, sulfonated	183873-19-0P	183873-20-3P	
183973-58-2P	183973-60-6P	183973-61-7P	184013-80-7P
(preparation of water-soluble fluorescent hybrid phthalocyanine derivs. for immunoassays)			
IT 19333-15-4P, Silicon phthalocyanine dihydroxide	37623-03-3P,		
1,4-Diphenyl-2,3-naphthalenedicarbonitrile	41345-70-4P,		
3-(Acetylthio)propanoic acid	52319-97-8P, 5-tert-Butyl-1,3-		
diiminoisoindoline	63405-81-2P, 5,6-Dichloro-1,3-		
diiminoisoindoline	92396-90-2P, Silicon naphthalocyanine		
dihydroxide	97241-14-0P, 1,3-Diiminoisoindoline-5,6-		
dicarbonitrile	111305-19-2P, 4,5,6,7-Tetrafluoro-1,3-		
diiminoisoindoline	121668-81-3P 163968-99-8P,		
4,9-Diethoxy-1,3-diiminobenz[f]isoindoline	163969-16-2P		
163969-17-3P	163969-19-5P, Dimethyl-7-octenylsilanol		
163969-20-8P, 4,9-Dibutoxy-1,3-diiminobenz[f]isoindoline			
163969-21-9P, 1,3-Diimino-4,9-diphenylbenz[f]isoindoline			
163969-23-1P, 6,7-Dibromo-1,3-diiminobenz[f]isoindoline			
183872-52-8DP, sulfonated	183872-52-8P 183872-53-9P		
183872-54-0P	183872-60-8P 183872-64-2P 183872-67-5P		
183872-68-6P, 4,7-Diethoxy-1,3-diiminoisoindoline	183872-69-7P		
183872-70-0P	183872-73-3P 183872-75-5P 183872-78-8P		
183872-80-2P	183872-83-5P 183872-91-5P 183872-93-7P		
183873-01-0P	183873-09-8DP, sulfonated 183873-16-7DP,		
sulfonated	183873-18-9P 183973-59-3P		
(preparation of water-soluble fluorescent hybrid phthalocyanine derivs. for immunoassays)			
IT 68-26-8, all-trans-Retinol	75-78-5 76-86-8,		
Chlorotriphenylsilane	79-08-3, Bromoacetic acid	107-96-0	
108-30-5, reactions	108-95-2, Phenol, reactions	111-87-5, 1-	
Octanol, reactions	140-66-9 597-52-4,		
Triethylsilanol	712-74-3, 1,2,4,5-		
Benzenetetracarbonitrile	1585-90-6, N-(2-Hydroxyethyl)maleimide		
1719-58-0, Chlorodimethylvinylsilane	1835-65-0,		
Tetrafluorophthalonitrile	2466-76-4, 1-Acetylimidazole		
3468-11-9, 1,3-Diiminoisoindoline	3634-67-1,		
Chlorotrihexylsilane	3663-43-2, (4-Aminobutyl)methoxydimethylsilane		
4655-61-2, 1,4-Dimethoxy-2,3-naphthalenedicarbonitrile			
6038-19-3, Homocysteine thiolactone hydrochloride	6554-98-9,		
trans-4-Hydroxystilbene	9004-74-4 10264-67-2,		
3,6-Diethoxyphthalonitrile	17196-12-2, Chlorodimethyl-7-		
octenylsilane	18156-15-5, Chloro(3-cyanopropyl)dimethylsilane		
18162-48-6, tert-Butylchlorodimethylsilane	18419-53-9,		
Chlorodiphenylvinylsilane	18643-08-8,		
Chlorodimethyloctadecylsilane	19333-10-9, Silicon phthalocyanine		
dichloride	20082-71-7, Chlorodimethyl(pentafluorophenyl)silane		
25322-68-3 26857-61-4	32703-80-3, 4-tert-Butylphthalonitrile		
36360-42-6, 3,6-Diphenylphthalonitrile	53749-38-5,		
(10-Carbomethoxydecyl)chlorodimethylsilane	74815-81-9,		
2,3-Dibromo-6,7-dicyanonaphthalene	92396-90-2D, sulfonated,		
compound with triethylamine	92396-91-3, Silicon naphthalocyanine		

dichloride 102488-47-1, Chlorodimethyl(3,3,4,4,5,5,6,6,7,7,8,8,8-
tridecafluoroctyl)silane 116453-89-5, 1,4-Dibutoxynaphthalene-
2,3-dicarbonitrile 116453-91-9, 1,4-Diethoxy-2,3-
naphthalenedicarbonitrile 139152-08-2, 4,5-
Dichlorophthalonitrile 163969-13-9 163969-22-0 183873-12-3
184047-30-1

(preparation of water-soluble fluorescent hybrid
phthalocyanine derivs. for immunoassays)

IT 57-27-2, Morphine, uses
(sulfonated hybrid phthalocyanine derivative conjugates;
preparation of water-soluble fluorescent hybrid phthalocyanine
derivs. as dye systems for immunoassays)

L15 ANSWER 22 OF 37 HCPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1996:175688 HCPLUS

DOCUMENT NUMBER: 124:203399

TITLE: Preparation of diphenylsiloxane-
dimethylsiloxane copolymers of narrow
molecular weight distribution

INVENTOR(S): Okawa, Tadashi

PATENT ASSIGNEE(S): Dow Corning Toray Silicone Co., Ltd., Japan

SOURCE: Eur. Pat. Appl., 9 pp.

CODEN: EPXXDW

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

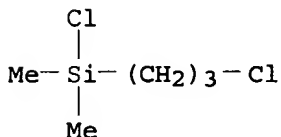
PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
EP 693521	A1	19960124	EP 1995-111454	1995 0720
R: DE, FR, GB			<--	
JP 08034855	A2	19960206	JP 1994-191075	1994 0721
			<--	
JP 3453430	B2	20031006		
US 5567790	A	19961022	US 1995-503601	1995 0714
			<--	
PRIORITY APPLN. INFO.:			JP 1994-191075	A
				1994 0721
			<--	

AB The title method comprises (I) polymerizing (A) a mixture of hexamethylcyclotrisiloxane and hexaphenylcyclotrisiloxane, using as a polymerization initiator (B) a Li metal salt of organosilane or polyorganosiloxane, optionally, in the presence of (C) a mol.-weight regulator selected from H₂O or a silanol compound, the polymerization taking place in the presence of both (D) a nitrile compound for restraining side reactions and (E) an active H-free polar solvent; and (II) terminating the polymerization reaction product from step (I) with a neutralizing agent selected from an acid or an organohalosilane. Dimethylsiloxane-diphenylsiloxane copolymer was made using Li adduct of siloxane diol oligomer, acetonitrile

inhibitor of side reactions, acetic acid chain terminator, and DMF solvent, and having number-average mol. weight 4870 and mol. weight distribution 1.15.

IT 10605-40-0
 (chain terminator via Li silanolate neutralization;
 nitrile side reaction inhibitor in preparation of
 diphenylsiloxane-dimethylsiloxane copolymers of narrow mol. weight
 distribution)

RN 10605-40-0 HCPLUS
 CN Silane, chloro(3-chloropropyl)dimethyl- (6CI, 7CI, 8CI, 9CI) (CA
 INDEX NAME)



IC ICM C08G077-08
 ICS C08G077-06
 CC 35-7 (Chemistry of Synthetic High Polymers)
 IT Polymerization
 (nitrile side reaction inhibitor in preparation of
 diphenylsiloxane-dimethylsiloxane copolymers of narrow mol. weight
 distribution)

IT Siloxanes and Silicones, preparation
 (di-Me, di-Ph, nitrile side reaction inhibitor in prepn
 . of diphenylsiloxane-dimethylsiloxane copolymers of narrow
 mol. weight distribution)

IT 64-19-7, Acetic acid, uses 79-09-4, Propionic acid, uses
 79-10-7, Acrylic acid, uses 124-38-9, Carbon dioxide, uses
 1066-35-9, Dimethylchlorosilane 1719-58-0,
 Dimethylvinylchlorosilane 7647-01-0, Hydrochloric acid, uses
 7664-93-9, Sulfuric acid, uses 10605-40-0 24636-31-5,
 Methacryloxypropyldimethylchlorosilane
 (chain terminator via Li silanolate neutralization;
 nitrile side reaction inhibitor in preparation of
 diphenylsiloxane-dimethylsiloxane copolymers of narrow mol. weight
 distribution)

IT 109-72-8D, n-Butyllithium, reaction product with
 siloxane oligomer 17574-46-8 58556-58-4 106211-26-1
 174614-54-1 174614-55-2 174614-56-3 174614-57-4
 174614-58-5 174614-59-6 174614-60-9
 (nitrile side reaction inhibitor in preparation of
 diphenylsiloxane-dimethylsiloxane copolymers of narrow mol. weight
 distribution)

IT 29300-68-3DP, Hexamethylcyclotrisiloxane-
 hexaphenylcyclotrisiloxane copolymer, hydroxy group-terminated
 (nitrile side reaction inhibitor in preparation of
 diphenylsiloxane-dimethylsiloxane copolymers of narrow mol. weight
 distribution)

IT 75-05-8, Acetonitrile, uses 78-82-0, Isobutyronitrile
 107-12-0, Propionitrile 109-74-0, Butyronitrile 110-59-8,
 Valeronitrile 110-61-2, Succinonitrile 140-29-4,
 α -Tolunitrile
 (nitrile side reaction inhibitor in preparation of
 diphenylsiloxane-dimethylsiloxane copolymers of narrow mol. weight
 distribution)

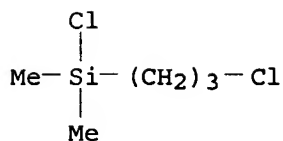
L15 ANSWER 23 OF 37 HCAPLUS COPYRIGHT 2006 ACS on STN
 ACCESSION NUMBER: 1996:175687 HCAPLUS
 DOCUMENT NUMBER: 124:203398
 TITLE: Preparation of siloxanes with
 minimal amount of low molecular weight
 organosiloxane
 INVENTOR(S): Manzouji, Ryuko; Okawa, Tadashi
 PATENT ASSIGNEE(S): Dow Corning Toray Silicone Co., Ltd., Japan
 SOURCE: Eur. Pat. Appl., 9 pp.
 CODEN: EPXXDW
 DOCUMENT TYPE: Patent
 LANGUAGE: English
 FAMILY ACC. NUM. COUNT: 1
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
EP 693520	A1	19960124	EP 1995-111453	1995 0720
EP 693520 R: DE, FR, GB JP 08034856	B1	20000823	<--	
	A2	19960206	JP 1994-191076	1994 0721
JP 3453431 US 5567789	B2	20031006	<--	
	A	19961022	US 1995-502447	1995 0714
PRIORITY APPLN. INFO.:			JP 1994-191076	A
				1994 0721
			<--	

AB The title method comprises (I) polymerizing (A) a mixture of cyclic trisiloxane, using as a polymerization initiator (B) a Li metal salt of organosilane or polyorganosiloxane, optionally, in the presence of (C) a mol.-weight regulator selected from H₂O or a **silanol** compound, the polymerization taking place in the presence of both (D) a nitrile compound for restraining side reactions and (E) an active H-free polar solvent; and (II) terminating the polymerization reaction product from step (I) with a neutralizing agent selected from an acid or an organohalosilane. Polydimethylsiloxane was made using Li adduct of siloxane diol oligomer, acetonitrile inhibitor of side reactions, acetic acid chain terminator, and DMF solvent, and having number-average mol. weight 29,411, mol. weight distribution 1.05, and dimethylsiloxane (d.p. ≤25) content 660 ppm.

IT 10605-40-0
 (chain terminator via Li **silanolate** neutralization;
 nitrile side reaction inhibitor in **preparation** of
 polydimethylsiloxane containing low amount siloxane oligomer)

RN 10605-40-0 HCAPLUS
 CN Silane, chloro(3-chloropropyl)dimethyl- (6CI, 7CI, 8CI, 9CI) (CA INDEX NAME)



IC ICM C08G077-08
 ICS C08G077-06
 CC 35-7 (Chemistry of Synthetic High Polymers)
 IT Polymerization
 (nitrile side reaction inhibitor in preparation of
 polydimethylsiloxane containing low amount siloxane oligomer)
 IT Siloxanes and Silicones, preparation
 (di-Me, di-Ph, nitrile side reaction inhibitor in prepn
 . of polydimethylsiloxane containing low amount siloxane oligomer)
 IT 64-19-7, Acetic acid, uses 79-09-4, Propionic acid, uses
 79-10-7, Acrylic acid, uses 124-38-9, Carbon dioxide, uses
 1066-35-9, Dimethylchlorosilane 1719-58-0,
 Dimethylvinylchlorosilane 7647-01-0, Hydrochloric acid, uses
 7664-93-9, Sulfuric acid, uses 10605-40-0 24636-31-5,
 Methacryloxypropyldimethylchlorosilane
 (chain terminator via Li silanolate neutralization;
 nitrile side reaction inhibitor in preparation of
 polydimethylsiloxane containing low amount siloxane oligomer)
 IT 109-72-8D, n-Butyllithium, reaction product with
 siloxane oligomer 17574-46-8 58556-58-4 106211-26-1
 174614-54-1 174614-55-2 174614-56-3 174614-57-4
 174614-58-5 174614-59-6 174614-60-9
 (nitrile side reaction inhibitor in preparation of
 polydimethylsiloxane containing low amount siloxane oligomer)
 IT 25084-99-5DP, Hexamethylcyclotrisiloxane homopolymer, hydroxy
 group-terminated 31692-79-2P
 (nitrile side reaction inhibitor in preparation of
 polydimethylsiloxane containing low amount siloxane oligomer)
 IT 75-05-8, Acetonitrile, uses 78-82-0, Isobutyronitrile
 107-12-0, Propionitrile 109-74-0, Butyronitrile 110-59-8,
 Valeronitrile 110-61-2, Succinonitrile 140-29-4,
 α -Tolunitrile
 (nitrile side reaction inhibitor in preparation of
 polydimethylsiloxane containing low amount siloxane oligomer)

L15 ANSWER 24 OF 37 HCPLUS COPYRIGHT 2006 ACS on STN
 ACCESSION NUMBER: 1994:186764 HCPLUS
 DOCUMENT NUMBER: 120:186764
 TITLE: Shielded stationary phases for liquid
 chromatography or extraction of mixtures containing
 proteins and small analytes
 INVENTOR(S): Feibusch, Binyamin; Gisch, Daryl J.
 PATENT ASSIGNEE(S): S.A.C. Corp., USA
 SOURCE: U.S., 19 pp. Cont. of U.S. Ser. No. 557,333,
 abandoned.
 CODEN: USXXAM
 DOCUMENT TYPE: Patent
 LANGUAGE: English
 FAMILY ACC. NUM. COUNT: 1
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
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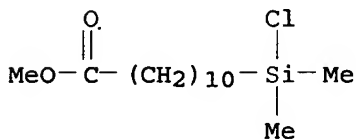
US 5277813	A	19940111	US 1992-988610	
				1992
				1210
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PRIORITY APPLN. INFO.:			US 1988-208200	B2
				1988
				0617
<--				
			US 1990-557333	B1
				1990
				0723
<--				

AB Novel packing materials are provided for liquid chromatog. and/or solid-phase extraction columns which will allow direct injection of biol. fluids for separation of small analytes from protein-containing mixts. These packing materials have a hydrophilic exterior layer and a hydrophobic, charged, or otherwise selective portion that forms an underlayer or is embedded in the hydrophilic layer. During a chromatog. process, large water-soluble biopolymers will be in contact with the hydrophilic outer layer and be shielded from interacting with the underlayer or embedded portion and elute unretained. Small analytes, on the other hand, can be fully partitioned throughout the exterior and interior layers and are retained by hydrophobic or electrostatic interactions. Silica- and silica gel-bonded phases were prepared [e.g., N,N-bis(2'-methoxyethyl)-11-(triethoxysilyl)undecylamine was prepared and bonded to silica gel] and used in the direct analyses of drugs in plasma or serum samples.

IT 53749-38-5
(reaction of, in preparation of shielded stationary phase for HPLC anal. of drugs in human blood serum)

RN 53749-38-5 HCAPLUS

CN Undecanoic acid, 11-(chlorodimethylsilyl)-, methyl ester (9CI)
(CA INDEX NAME)



IC ICM B01D015-08

INCL 210502100

CC 9-3 (Biochemical Methods)

Section cross-reference(s): 1, 4, 29, 66

IT 50-06-6, analysis 50-47-5, Desipramine 51-06-9, Procainamide 56-54-2, Quinidine 56-75-7, Chloramphenicol 58-08-2, analysis 58-55-9, Theophylline, analysis 58-93-5, Hydrochlorothiazide 65-85-0, Benzoic acid, analysis 66-22-8, Uracil, analysis 69-72-7, analysis 76-57-3, Codeine 103-90-2, Acetaminophen 298-46-4, Carbamazepine 525-66-6, Propranolol 738-70-5 15687-27-1, Ibuprofen
(determination of, in human blood serum by HPLC using shielded stationary phases)

IT 111-26-2, Hexylamine 124-09-4, 1,6-Hexanediamine, uses 2052-49-5, Tetrabutylammonium hydroxide

(in preparation of shielded stationary phase for HPLC anal. of drugs in human blood serum)

IT 116047-42-8P, 11-Triethoxysilylundecanal
(preparation and reaction of, in preparation of shielded stationary phase for HPLC anal. of drugs in human blood serum)

IT 7631-86-9DP, Silica, reaction products with hydrophobic compds. having polar functionality 40762-31-0DP, reaction products with Supelcosil silica and propane sultone 95752-11-7DP, Supelcosil DB, reaction products with hydrophobic compds. having polar functionality 153632-55-4DP, reaction products with silica gel 153632-57-6DP, reaction products with silica gel
(preparation of, as shielded stationary phase for HPLC anal. of drugs in human blood serum)

IT 108-30-5, Succinic anhydride, analysis 111-95-5, Bis(2-methoxyethyl)amine 1120-71-4, 1,3-Propane sultone 2489-52-3, 3-Fluorosulfonylbenzenesulfonyl chloride 2530-86-1 13822-56-5, 3-Aminopropyltrimethoxysilane 20493-87-2, N-(2-Aminoethyl)-3-aminopropyltrimethylsilane 35141-36-7, N-(3-Trimethoxysilylpropyl)trimethylammonium chloride 40762-30-9 53749-38-5 71245-36-8, Hypol FHP 2000 72876-91-6 105053-76-7, AZ-CUP MC 147366-30-1, N-(3-Trimethoxysilylpropyl)tributylammonium bromide 153632-56-5, N-Hydroxysuccinimido 11-(triethoxysilyl)undecanoate
(reaction of, in preparation of shielded stationary phase for HPLC anal. of drugs in human blood serum)

IT 112-45-8, 10-Undecenal
(reaction of, with triethoxysilane, in preparation of shielded stationary phase for HPLC anal. of drugs in human blood serum)

IT 998-30-1, Triethoxysilane
(reaction of, with undecenal, in preparation of shielded stationary phase for HPLC anal. of drugs in human blood serum)

L15 ANSWER 25 OF 37 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1992:583782 HCAPLUS

DOCUMENT NUMBER: 117:183782

TITLE: Preparation of well-engineered thin molecular layers on semiconductor-based transducers

AUTHOR(S): Jaffrezic-Renault, N.; Martelet, C.

CORPORATE SOURCE: Lab. Physicochem. Interfaces, Ec. Cent. Lyon, Ecully, F-69131, Fr.

SOURCE: Sensors and Actuators, A: Physical (1992), A32(1-3), 307-12

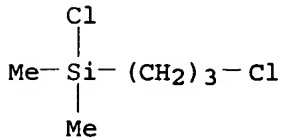
CODEN: SAAPEB; ISSN: 0924-4247

DOCUMENT TYPE: Journal; General Review

LANGUAGE: English

AB Chemical and biochem. sensors, based on semiconductor substrates such as ISFETs, enzymic FETs (ENFETs) or capacitive immunosensors, need as a sensing part a structure exhibiting good mol. or ionic recognition properties associated with a stability allowing a long lifetime. In order to achieve such requirements the authors developed direct chemical grafting techniques onto active or passive electronic components such as FET or electrolyte oxide semiconductor (EOS) capacitors. Examples of ISFETs (Ag+, Ca2+, NO3-), ENFETs (urea, glucose) and immunosensors (IgG, IgE) are given to emphasize the advantages of this direct coupling process of a monomol. layer onto a semiconductor-based

transducer.
 IT 10605-40-0
 (reaction of, with silanol groups on silicon surface,
 for ionophore fixation in preparation of sensors)
 RN 10605-40-0 HCPLUS
 CN Silane, chloro(3-chloropropyl)dimethyl- (6CI, 7CI, 8CI, 9CI) (CA
 INDEX NAME)



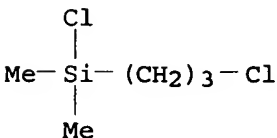
CC 79-2 (Inorganic Analytical Chemistry)
 Section cross-reference(s): 9, 15, 76
 ST thin mol layer prepn semiconductor transducer; review
 thin mol layer semiconductor sensor; grafting technique
 semiconductor transducer surface; FET sensor monomol layer film
 prepн; sensor semiconductive thin mol layer prepн
 ; biosensor semiconductive monomol layer film prepн;
 immunosensor semiconductive monomol layer film prepн
 IT Immunoassay
 (grafting technique for preparation of well-engineered
 thin mol. layers on capacitor for)
 IT Semiconductor devices
 (grafting technique for preparation of well-engineered
 thin mol. layers on, for anal.)
 IT Sensors
 (semiconductor-based, grafting technique for preparation
 of well-engineered thin mol. layers for, for anal.)
 IT Immunoglobulins
 (E, determination of, grafting technique for preparation of
 well-engineered thin mol. layers on capacitor for)
 IT Immunoglobulins
 (G, determination of, grafting technique for preparation of
 well-engineered thin mol. layers on capacitor for)
 IT Transistors
 (field-effect, grafting technique for preparation of
 well-engineered thin mol. layers on, for anal.)
 IT 50-99-7, Glucose, analysis 57-13-6, Urea, analysis 7440-22-4,
 Silver, analysis 7440-70-2, Calcium, analysis 14797-55-8,
 Nitrate, analysis 16637-16-4, Uranyl ion(2+)
 (determination of, grafting technique for preparation of
 well-engineered thin mol. layers on FET for)
 IT 111-30-8, Pentanedral
 (reaction of, with silanized silicon surface, for protein
 fixation in preparation of calcium sensor)
 IT 141630-76-4 143982-95-0
 (reaction of, with silanized silicon surface, in prepн
 . of calcium sensor)
 IT 143982-94-9
 (reaction of, with silanized silicon surface, in prepн
 . of uranyl sensor)
 IT 3663-43-2
 (reaction of, with silanol groups on silicon surface,
 for fixation of biospecific mols. in preparation of
 sensors)

IT 10605-40-0
 (reaction of, with silanol groups on silicon surface,
 for ionophore fixation in preparation of sensors)
 IT 22705-32-4D, derivs.
 (reaction of, with silanol groups on silicon surface,
 in preparation of sensors)

L15 ANSWER 26 OF 37 HCAPLUS COPYRIGHT 2006 ACS on STN
 ACCESSION NUMBER: 1992:476362 HCAPLUS
 DOCUMENT NUMBER: 117:76362
 TITLE: Novel silicones for transdermal therapeutic
 system. III. Preparation of
 pyridinio- or ammonio-terminated
 polydimethylsiloxanes and the evaluation as
 transdermal penetration enhancers
 AUTHOR(S): Aoyagi, Takao; Nakamura, Tomoko; Yabuchi,
 Yuichi; Nagase, Yu
 CORPORATE SOURCE: Sagami Chem. Res. Cent., Sagamihara, 229,
 Japan
 SOURCE: Polymer Journal (Tokyo, Japan) (1992
), 24(6), 545-53
 CODEN: POLJB8; ISSN: 0032-3896
 DOCUMENT TYPE: Journal
 LANGUAGE: English

AB α -[3-(1-Pyridinio)propyl]polydimethylsiloxane iodide
 (PDMS-Py+I- and α -[3-(N,N-dimethylethylammonio)propyl]polydimethylsiloxane iodide (PDMS-Am+I-) were prepared from PDMS containing iodopropyl group at the chain end. The prepolymer was synthesized by a ring-opening polymerization of hexamethylcyclotrisiloxane (D3) initiated with lithium trimethylsilanolate followed by the termination with 3-chloropropyltrimethylchlorosilane and the halogen substitution with NaI. All the polymers effectively enhanced the drug penetration through the skin and the permeation coeffs. were about 2-6 times as much as that without enhancer. It was revealed from the detailed anal. of the permeation profile that the permeation and partition coeffs. increased in parallel, with increasing of the average degree of polymerization, while the diffusion coeffs. were unchanged.

IT 10605-40-0DP, 3-Chloropropyltrimethylchlorosilane, reaction
 products with hexamethylcyclotrisiloxane
 (preparation and chloride substitution reaction of)
 RN 10605-40-0 HCAPLUS
 CN Silane, chloro(3-chloropropyl)dimethyl- (6CI, 7CI, 8CI, 9CI) (CA
 INDEX NAME)



CC 63-5 (Pharmaceuticals)
 Section cross-reference(s): 35
 IT Siloxanes and Silicones, biological studies
 (di-Me, quaternary ammonium group-containing, preparation and
 evaluation as transdermal penetration enhancer of)
 IT Pharmaceutical dosage forms

(transdermal, penetration enhancers for, quaternary ammonium-containing polydimethylsiloxanes as, preparation and evaluation of)

IT 10605-40-0DP, 3-Chloropropyldimethylchlorosilane, reaction products with hexamethylcyclotrisiloxane
(preparation and chloride substitution reaction of)

IT 110-86-1DP, Pyridine, reaction products with polydimethylsiloxane 598-56-1DP, N,N-Dimethylethylamine, reaction products with polydimethylsiloxane 9016-00-6DP, Poly[oxy(dimethylsilylene)], ammonio- and pyridinio-terminated
(preparation and evaluation as transdermal penetration enhancer of)

IT 25084-99-5DP, ammonio- and pyridinio-terminated
(preparation and evaluation as transdermal penetration enhancer of)

L15 ANSWER 27 OF 37 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1992:210721 HCAPLUS

DOCUMENT NUMBER: 116:210721

TITLE: Fluorescent porphyrin and fluorescent phthalocyanine-polyethylene glycol, -polyol, and saccharide derivatives as fluorescent probes

INVENTOR(S): Arrhenius, Peter Olof Gustaf

PATENT ASSIGNEE(S): Diatron Corp., USA

SOURCE: PCT Int. Appl., 43 pp.
CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 9

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 9118006	A1	19911128	WO 1991-US3424	1991 0515 ---
CA 2082936	AA	19911116	CA 1991-2082936	1991 0515 ---
CA 2082936	C	20030923		1991 0515 ---
EP 529002	A1	19930303	EP 1991-912121	1991 0515 ---
EP 529002	B1	20011004		1991 0515 ---
JP 05508015	R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE T2	19931111	JP 1991-511180	1991 0515 ---
JP 3224538	B2	20011029		1991 0515 ---
US 5403928	A	19950404	US 1991-701449	1991

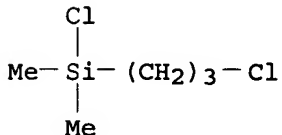
			0515
AT 206395	E	20011015	<-- AT 1991-912121
			1991 0515
ES 2163393	T3	20020201	<-- ES 1991-912121
			1991 0515
PRIORITY APPLN. INFO.:			<-- US 1990-523601 A 1990 0515
			<-- US 1991-701449 1991 0515
			<-- WO 1991-US3424 W 1991 0515
			<--

AB Marker components are prepared which are compatible with aqueous solns., exhibit favorable fluorescence properties, and exhibit decreased nonspecific binding to macromols. in solution. These markers are useful for e.g. fluorescence immunoassays. A digoxigenin probe was prepared by conjugating 2,3-dicarboxyphthalocyaninato-bis[3-(1H-imidazol-1-ylcarbonyl) aminopropyldimethylsilanolate]silicon (preparation given) with amine-terminated PEG (preparation given). The product was further conjugated with 3-dl-aminodigoxigenin. The product gave an immunospecific reaction with a specific digoxin antibody.

IT 10605-40-0
(reaction of, in conjugate fluorescent probe preparation)

RN 10605-40-0 HCAPLUS

CN Silane, chloro(3-chloropropyl)dimethyl- (6CI, 7CI, 8CI, 9CI) (CA INDEX NAME)



IC C07J043-00; C07D209-56; C07D487-22; C07H023-00

CC 9-5 (Biochemical Methods)

Section cross-reference(s): 15, 29

IT Alcohols, compounds

(polyhydric, conjugates, with fluorophores, for fluorescent probes)

IT 3468-11-9P 19333-15-4P 25322-68-3DP, amine-terminated
97241-14-0P 140871-08-5P 140871-09-6P 140871-10-9P
140889-30-1P 140890-47-7P

(preparation and reaction of, in conjugate fluorescent probe preparation)

IT 448-65-7DP, Deuteroporphyrin, mannitol esters 83830-83-5DP,
reaction products with fluorophore-PEG derivative conjugate

140871-07-4P 140871-09-6DP, amine-terminated PEG conjugates
 140888-76-2P 140889-30-1DP, amine-terminated PEG conjugates
 140890-46-6P 140907-65-9P 140935-72-4P
 (preparation of, for fluorescent probe)
 IT 69-65-8, Mannitol 91-15-6, Phthalonitrile 288-32-4, Imidazole, reactions 448-65-7, Deuteroporphyrin 10026-04-7, Silicon tetrachloride 10605-40-0 14459-29-1, Hematoporphyrin IX 15554-15-1, Hydroxyaluminum phthalocyanine 17070-70-1, 3-Isocyanatopropyldimethylchlorosilane 27879-07-8, Polyethyleneglycol monoethyl ether 83830-83-5 140890-48-8 (reaction of, in conjugate fluorescent probe preparation)
 IT 108-30-5, reactions (reaction of, with phthalocyanine derivative in conjugate fluorescent probe preparation)

L15 ANSWER 28 OF 37 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1992:173516 HCAPLUS
 DOCUMENT NUMBER: 116:173516
 TITLE: Supramolecular asymmetric induction with a NADH model reagent grafted on silica
 Losset, D.; Dupas, G.; Duflos, J.; Bourguignon, J.; Queguiner, G.
 CORPORATE SOURCE: Lab. Chim. Org. Fine Heterocyclique, INSA, Mont-Saint Aigman, 76131, Fr.
 SOURCE: Bulletin de la Societe Chimique de France (1991), (Sept.-Oct.), 721-9
 CODEN: BSCFAS; ISSN: 0037-8968

DOCUMENT TYPE: Journal
 LANGUAGE: French

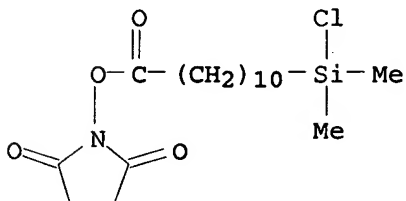
OTHER SOURCE(S): CASREACT 116:173516

AB A model of NADH issued from a thieno[2,3-b]dihydropyridine derivative has been grafted onto a silica matrix bearing on another part a chiral auxiliary. Two strategies were implemented to obtain the corresponding reagents. In the first case, the reagent and the auxiliary were grafted to the silica by means of two different arms. In the second case, the reagent and the auxiliary were linked to two arms which are brought together before being linked to the silica matrix. The reagents thus obtained were involved in the reduction of Me phenylglyoxylate and enantiomeric excesses of 20 and 35% were obtained.

IT 139764-32-2P (preparation and reaction of, with silica)

RN 139764-32-2 HCAPLUS

CN 2,5-Pyrrolidinedione, 1-[[11-(chlorodimethylsilyl)-1-oxoundecyl]oxy]- (9CI) (CA INDEX NAME)



CC 23-7 (Aliphatic Compounds)
 Section cross-reference(s): 22

IT 139764-37-7P (preparation and condensation of, with

dimethylphenylalaninol)
 IT 7766-49-6P 110661-49-9P 139764-38-8P
 (preparation and hydrosilylation of)
 IT 139764-35-5DP, reaction products with silica and
 (chlorodimethylsilyl)iododecane 139764-40-2DP, silica-bound
 (preparation and reaction of, with
 carbamoylthienopyridine)
 IT 139764-34-4DP, reaction products with silica and
 [[[chlorosilyl]decyl]carbonyloxy]succinimide 139764-43-5DP,
 reaction products with silica and
 hydroxy(iodoundecyl)silane
 (preparation and reaction of, with dimethylphenylalaninol)
 IT 139764-32-2P 139764-33-3P 139764-39-9P
 (preparation and reaction of, with silica)
 IT 139764-36-6DP, reaction products with silica and
 hydroxy[(alanylcarbonyl)oxy]decylsilane 139764-41-3DP,
 silica-bound
 (preparation and reduction by, of glyoxylate)
 IT 140168-98-5DP, silica-bound
 (preparation and reduction of)
 IT 140168-99-6DP, reaction products with silica and
 hydroxy[[[(dimethylamino)propyl]oxy]carbonyl]decyl]silane
 (preparation and regioselective reduction of)
 IT 15206-55-0P 20698-91-3P 21210-43-5P
 (preparation of)
 IT 1931-60-8
 (sequential reactions of, with bromoundecanol and
 hydroxydodecanoic acid)

L15 ANSWER 29 OF 37 HCPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1991:164493 HCPLUS
 DOCUMENT NUMBER: 114:164493
 TITLE: Preparation of
 (chlorodimethylsilyl)carboxylic acid
 trialkylsilyl esters
 INVENTOR(S): Tezuka, Yasushi; Imai, Kiyokazu
 PATENT ASSIGNEE(S): Toshiba Silicone Co., Ltd., Japan
 SOURCE: Jpn. Kokai Tokkyo Koho, 9 pp.
 CODEN: JKXXAF
 DOCUMENT TYPE: Patent
 LANGUAGE: Japanese
 FAMILY ACC. NUM. COUNT: 1
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
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JP 02282389	A2	19901119	JP 1989-102358	1989 0421

PRIORITY APPLN. INFO.: <--
 JP 1989-102358
 1989
0421

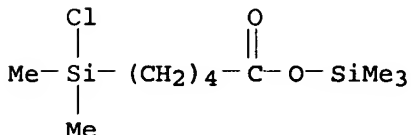
OTHER SOURCE(S): MARPAT 114:164493
 AB ClSiMe2ZCO2SiMe2R (I: R = C1-4 alkyl; Z = C1-6 hydrocarbylene
 containing no aliphatic unsatd. bond), useful as synthetic intermediates
 and coupling agents, were prepared Me3SiCl was gradually
 added to a mixture of CH2:CHCH2CO2H, Et2O, and Et3N at 0° and

the reaction mixture was refluxed for 2 h to give 65% $\text{CH}_2:\text{CHCH}_2\text{CO}_2\text{SiMe}_3$ which was treated with HSiClMe_2 and Pt/C while gradually heating to 45° and refluxing the mixture for 2 h to give 53% I [R = Me, Z = $(\text{CH}_2)_3$].

IT 130200-19-OP
(preparation of, as coupling agent)

RN 130200-19-0 HCAPLUS

CN Pentanoic acid, 5-(chlorodimethylsilyl)-, trimethylsilyl ester
(9CI) (CA INDEX NAME)



IC ICM C07F007-18
CC 29-6 (Organometallic and Organometalloidal Compounds)

IT Carboxylic acids, esters

(silyl, esters, with trialkylsilanols, prepns. of, as coupling agents)

IT 13688-54-5P 23523-56-0P

(preparation and hydrosilylation of, with dimethylchlorosilane)

IT 130200-18-9P 130200-19-0P

(preparation of, as coupling agent)

115 ANSWER 30 OF 37 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1991:143533 HCAPLUS

ACCESSION NUMBER: 1991.14353
DOCUMENT NUMBER: 114:143533

DOCUMENT NUMBER: 114.143333
TITLE: Fast atom bombardment mass spectrometry of some alkoxy- and chlorosilanes

AUTHOR(S): Kallury, Krishna M. R.; Krull, Ulrich J.; Thompson, Michael

CORPORATE SOURCE: Thompson, Michael
Dep. Chem., Univ. Toronto, Toronto, ON, M5S
1A1, Can.

SOURCE: *Organic Mass Spectrometry (1991)*, 26 (2), 81-4

CODEN: ORMSBG; ISSN: 0030-493X

DOCUMENT TYPE: Journal

LANGUAGE: English

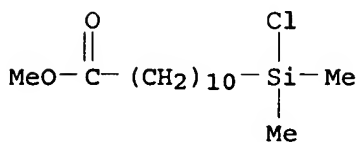
AB The pos. and neg. FAB mass spectra of a series of alkoxy- and chlorosilanes $Xm(CH_3)_{3-m}Si(CH_2)_nR$ [$m = 1$ or 3 , $n = 3, 10$ or 17 , $X = Cl$ or OMe or OEt , $R = Me, NH_2$, glycidoxy, CO_2Me , $NHCO(CH_2)_7CO_2Me$ or $NHCO(CH_2)_{10}CH_2OAc$] were recorded in NBA and NPOE matrixes. The chlorosilanes underwent rapid hydrolysis into silanols which condense to form siloxanes, the process being complete in NBA and partial in NPOE, yielding siloxane-based fragment ions in the pos. spectra and silyloxyanions in the neg. spectra. The alkoxy silanes were more resistant to hydrolysis, affording abundant $[MH-HX]^+$ ions ($X = OMe$ or OEt) in their pos. FAB spectra and moderate to high intensity $[M-H]^-$ ions in the neg. mode, the latter undergoing characteristic sequential loss of C_2H_4 , $EtOH$ and C_2H_4 . Significant variations were observed in the pos. spectra of all the silanes with change of matrix.

IT 53749-38-5

(pos. FAB mass spectrum of)

RN 53749-38-5 HCAPLUS

CN Undecanoic acid, 11-(chlorodimethylsilyl)-, methyl ester (9CI)
(CA INDEX NAME)

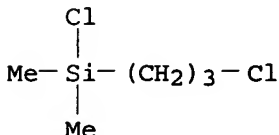


CC 29-6 (Organometallic and Organometalloidal Compounds)
IT 112-04-9 919-30-2 2530-83-8 18306-79-1 18643-08-8
53749-38-5 58160-70-6 132933-18-7 132933-19-8
(pos. FAB mass spectrum of)

L15 ANSWER 31 OF 37 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1989:632954 HCAPLUS
DOCUMENT NUMBER: 111:232954
TITLE: The modification of reactivity at a silicon center by a remote phosphorus group
AUTHOR(S): Kowalski, Jozef; Chojnowski, Julian
CORPORATE SOURCE: Cent. Mol. Macromol. Stud., Pol. Acad. Sci., Lodz, 90-362, Pol.
SOURCE: Journal of Organometallic Chemistry (1988), 356(3), 285-95
CODEN: JORCAI; ISSN: 0022-328X
DOCUMENT TYPE: Journal
LANGUAGE: English
OTHER SOURCE(S): CASREACT 111:232954

AB The compds. $\text{X}(\text{CH}_2)_n\text{SiMe}_2(\text{OPh})$ [$\text{X} = \text{H}$, $n = 2, 3$; $\text{X} = \text{PPh}_2$, $n = 1, 2, 3$; $\text{X} = \text{P}(\text{O})\text{Ph}_2$, $n = 2, 3$; $\text{X} = \text{P}(\text{S})\text{Ph}_2$, $n = 1, 2, 3$] having Si and P bridged by C chains, have been synthesized. The kinetics of acid- and base-catalyzed solvolytic cleavage of the phenoxy group from these compds. in methanol have been investigated. The kinetic results obtained in the presence of bases can be interpreted in terms of polar and steric effects alone, but there was an unexpected enhancement of the reactivity in the case of the P:O-containing substrates in acidic media. The solvent kinetic isotope effects are best interpreted in terms of participation by the P:O group as a base rather than as a nucleophile attacking the Si center.

IT 10605-40-0
(phenoxy substitution of)
RN 10605-40-0 HCAPLUS
CN Silane, chloro(3-chloropropyl)dimethyl- (6CI, 7CI, 8CI, 9CI) (CA INDEX NAME)



CC 29-6 (Organometallic and Organometalloidal Compounds)
Section cross-reference(s): 22, 67
IT 4028-23-3 6917-76-6 10605-40-0 17477-29-1
(phenoxy substitution of)

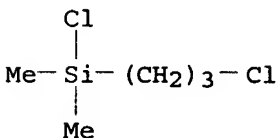
IT 124007-96-1P
 (preparation and reaction of, with chlorophosphines)
 IT 66998-68-3P
 (preparation and reaction of, with phosphines)
 IT 17876-92-5P 103676-01-3P 103676-03-5P 124007-89-2P
 124007-90-5P 124007-91-6P 124007-92-7P 124007-93-8P
 124007-94-9P 124007-95-0P
 (preparation and solvolysis of, kinetics of)

L15 ANSWER 32 OF 37 HCAPLUS COPYRIGHT 2006 ACS on STN
 ACCESSION NUMBER: 1989:574221 HCAPLUS
 DOCUMENT NUMBER: 111:174221
 TITLE: Synthesis of siloxyphosphines
 AUTHOR(S): Urbaniak, W.; Marciniec, B.
 CORPORATE SOURCE: Fac. Chem., A. Mickiewicz Univ., Poznan,
 60-780, Pol.
 SOURCE: Synthesis and Reactivity in Inorganic and
 Metal-Organic Chemistry (1988),
 18(7), 695-703
 CODEN: SRIMCN; ISSN: 0094-5714

DOCUMENT TYPE: Journal
 LANGUAGE: English
 OTHER SOURCE(S): CASREACT 111:174221

AB Siloxyphosphines $[\text{Ph}_2\text{P}(\text{CH}_2)_n\text{SiMe}_2]_2\text{O}$ ($n = 1, 3$) were prep'd
 by phosphidation of $\text{X}(\text{CH}_2)_n\text{SiMe}_2\text{OEt}$ ($\text{X} = \text{Br}$, $n = 1$; $\text{X} = \text{Cl}$, $n = 3$) with Ph_2PLi followed by hydrolysis. Free-radical addition of
 Ph_2PH to suitable vinylsiloxanes yielded $(\text{Ph}_2\text{PCH}_2\text{CH}_2\text{SiMe}_2)_2\text{O}$ and
 $(\text{Ph}_2\text{PCH}_2\text{CH}_2\text{SiMe}_2\text{O})_2\text{SiMeCH}_2\text{CH}_2\text{PPh}_2$.

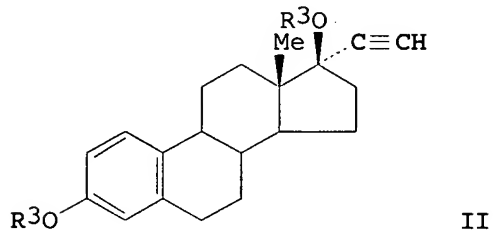
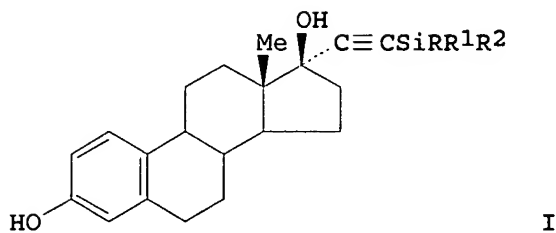
IT 10605-40-0
 (hydrolysis and ethanolysis of)
 RN 10605-40-0 HCAPLUS
 CN Silane, chloro(3-chloropropyl)dimethyl- (6CI, 7CI, 8CI, 9CI) (CA
 INDEX NAME)



CC 29-7 (Organometallic and Organometalloidal Compounds)
 IT 10605-40-0 16532-02-8, (Bromomethyl)chlorodimethylsilane
 (hydrolysis and ethanolysis of)
 IT 2351-13-5P 18132-72-4P
 (preparation and phosphidation of)
 IT 110547-70-1P, Bis(diphenylphosphinomethyl)tetramethyldisiloxane
 110547-71-2P 110547-72-3P 110547-73-4P
 (preparation of)
 IT 13508-63-9P 18156-50-8P
 (preparation, phosphidation, and subsequent hydrolysis of)

L15 ANSWER 33 OF 37 HCAPLUS COPYRIGHT 2006 ACS on STN
 ACCESSION NUMBER: 1987:138687 HCAPLUS
 DOCUMENT NUMBER: 106:138687
 TITLE: Steroidal silicon side-chain analogs as
 potential antifertility agents
 AUTHOR(S): Peters, Richard H.; Crowe, David F.; Tanabe,
 Masato; Avery, Mitchell A.; Chong, Wesley K.

CORPORATE SOURCE: M. Bio-Org. Chem. Lab., SRI Int., Menlo Park, CA, 94025, USA
 SOURCE: Journal of Medicinal Chemistry (1987), 30(4), 646-52
 DOCUMENT TYPE: CODEN: JMCMAR; ISSN: 0022-2623
 LANGUAGE: Journal
 OTHER SOURCE(S): English
 CASREACT 106:138687
 GI

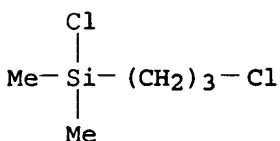


AB Ethynylestradiol Si analogs I [SiRR1R2 = SiMe3, SiEt3, SiPr3, SiEt3(CMe3), SiMe2Pr, etc.] were prepared by the reaction of ethynylestradiols II (R3 = H, THP) with ClSiRR1R2 in the presence of MeMg Br or BuLi followed by acid-catalyzed methanolysis. I exhibit high antifertility potency and markedly reduced estrogenic activity. The best compds. are I (R-R2 = Et; R = R1 = Me, R2 = CMe3), which show a separation of antifertility from estrogenic activity in rats. Structure-activity studies indicated a good correlation between biol. activities and calculated van der Waals vols. of R, R1, and R2.

IT 10605-40-0
 (silylation by, of ethynylestradiol)

RN 10605-40-0 HCAPLUS

CN Silane, chloro(3-chloropropyl)dimethyl- (6CI, 7CI, 8CI, 9CI) (CA INDEX NAME)



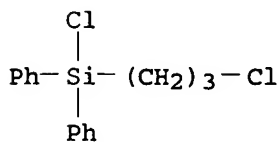
CC 32-3 (Steroids)
 Section cross-reference(s): 1
 ST ethynylestradiol silicon analog prepn antifertility;

estradiol ethynyl silicon analog
 IT 19-Norsteroids
 (silicon side-chain analogs, preparation and antifertility activities of)
 IT 597-49-9, 3-Ethyl-3-pentanol
 (chlorination of)
 IT 57-63-6DP, silyl derivs.
 (preparation and antifertility activities of)
 IT 50866-93-8P 50866-94-9P 50866-95-0P 50866-96-1P
 50866-97-2P 50866-98-3P 50866-99-4P 50867-00-0P
 50867-01-1P 50867-02-2P 50867-03-3P 50867-04-4P
 50867-05-5P 50867-06-6P 50867-07-7P 50867-08-8P
 50867-09-9P 50867-10-2P 50867-11-3P 50867-12-4P
 50938-72-2P 50938-73-3P 57099-89-5P 57099-90-8P
 57099-91-9P 57099-92-0P 57099-93-1P 57099-94-2P
 107149-27-9P 107149-28-0P 107149-29-1P 107149-30-4P
 107149-31-5P 107149-32-6P 107149-33-7P 107149-34-8P
 107149-35-9P 107149-36-0P 107149-37-1P 107149-38-2P
 107149-39-3P 107149-40-6P 107149-41-7P 107149-42-8P
 107149-43-9P 107149-44-0P 107149-45-1P 107149-46-2P
 107149-47-3P 107149-48-4P 107149-49-5P 107149-50-8P
 107149-51-9P 107149-52-0P 107149-53-1P 107173-94-4P
 107173-95-5P 107173-96-6P
 (preparation and antifertility activity of)
 IT 919-23-3P, 3,3-Diethyl-1-pentyne
 (preparation and reaction of, with estrone
 tetrahydropyranyl ether)
 IT 994-25-2P, 3-Chloro-3-ethylpentane
 (preparation and reaction of, with vinyl chloride)
 IT 75-77-4, Trimethylsilyl chloride, reactions 76-86-8,
 Triphenylsilyl chloride 768-33-2 994-30-9 995-04-0
 995-25-5 995-45-9, Tributylsilyl chloride 1000-50-6
 1481-41-0 1719-57-9 1833-31-4 3634-56-8 4028-23-3
 7787-82-8 10605-40-0 16532-02-8 17477-29-1
 17876-59-4 18148-37-3 18162-48-6 18163-33-2 18171-56-7
 18171-59-0 18279-72-6 18293-66-8 19923-52-5 57099-95-3
 60090-96-2 107149-54-2 107149-55-3 107149-56-4 107149-57-5
 (silylation by, of ethynylestradiol)

L15 ANSWER 34 OF 37 HCAPLUS COPYRIGHT 2006 ACS on STN
 ACCESSION NUMBER: 1980:42039 HCAPLUS
 DOCUMENT NUMBER: 92:42039
 TITLE: Sila drugs. 11. Diphenyl(3-piperidinopropyl)
 silanol, a sila analog of diphenidol
 Steiling, Lothar; Tacke, Reinhold; Wannagat,
 Ulrich
 AUTHOR(S): Inst. Anorg. Chem., Tech. Univ. Braunschweig,
 Braunschweig, D-3300, Fed. Rep. Ger.
 CORPORATE SOURCE: Liebigs Annalen der Chemie (1979),
 (10), 1554-9
 SOURCE: CODEN: LACHDL; ISSN: 0170-2041
 DOCUMENT TYPE: Journal
 LANGUAGE: German
 OTHER SOURCE(S): CASREACT 92:42039
 AB Successive Grignard reaction of $\text{Cl}_3\text{Si}(\text{CH}_2)_3\text{Cl}$ with PhMgCl gave
 $\text{Ph}_2\text{SiCl}(\text{CH}_2)_3\text{Cl}$, successive amine substitution of which with
 piperidine (QH) gave $\text{Ph}_2\text{SiQ}(\text{CH}_2)_3\text{Q}$ (I). Hydrolysis of I gave
 $\text{Ph}_2\text{Si}(\text{OH})(\text{CH}_2)_3\text{Q}$ (II), a sila analog of difenidol, which was
 quaternized with MeI. The LD₅₀ for II in the guinea pig was 101.6
 mg/kg. II was effective as an antiarrhythmic, anticholinergic,

histaminolytic, and muscle relaxant.

IT 2632-94-2P
 (preparation and amine substitution of, with piperidine)
 RN 2632-94-2 HCPLUS
 CN Silane, chloro(3-chloropropyl)diphenyl- (7CI, 8CI, 9CI) (CA INDEX NAME)



CC 29-6 (Organometallic and Organometalloidal Compounds)
 ST antihistamine piperidinopropyldiphenylsilanol;
 histaminolytic piperidinopropyldiphenylsilanol;
 anticholinergic piperidinopropyldiphenylsilanol;
 spasmolytic piperidinopropyldiphenylsilanol; muscle
 relaxant piperidinopropyldiphenylsilanol; antiarrhythmic
 piperidinopropyldiphenylsilanol; parasympatholytic
 piperidinopropyldiphenylsilanol; silanol
 piperidinopropyl diphenyl; difenidol sila analog
 IT Antiarrhythmics
 Antihistaminics
 Muscle relaxants and Spasmolytics
 Parasympatholytics
 (diphenyl(piperidinylpropyl)silanols)
 IT 2550-06-3P 3401-26-1P
 (preparation and Grignard reaction of, with chlorobenzene)
 IT 2632-94-2P 72315-20-9P
 (preparation and amine substitution of, with piperidine)
 IT 72315-21-0P
 (preparation and hydrolysis of)
 IT 72191-17-4P
 (preparation and quaternization of, with Me iodide)
 IT 72315-22-1P
 (preparation of)

L15 ANSWER 35 OF 37 HCPLUS COPYRIGHT 2006 ACS on STN
 ACCESSION NUMBER: 1973:442603 HCPLUS
 DOCUMENT NUMBER: 79:42603
 TITLE: Reaction of organosilicon alcohols
 and phenols with phosgene
 AUTHOR(S): Mironov, V. F.; Sheludyakov, V. D.;
 Khatuntsev, G. D.; Kozlikov, V. L.
 CORPORATE SOURCE: USSR
 SOURCE: Zurnal Obshchey Khimii (1973),
 43(3), 616-20
 CODEN: ZOKHA4; ISSN: 0044-460X
 DOCUMENT TYPE: Journal
 LANGUAGE: Russian
 AB COCl_2 reacted at 0-10° with $\text{O}(\text{SiMe}_2\text{ZOH})_2$ (I; Z = CH₂,
 $(\text{CH}_2)_3$, $\text{CH}_2\text{OCH}_2\text{CH}_2$) to form $\text{O}(\text{SiMe}_2\text{ZO}_2\text{CCl})_2$ (II), $\text{SiMe}_2(\text{ZCl})\text{Cl}$
 (III), and $\text{ClSiMe}_2\text{ZO}_2\text{CCl}$ (IV); the ratios were controlled by
 reactant ratios and the nature of Z. Thus, passing COCl_2 into
 $\text{O}(\text{SiMe}_2\text{CH}_2\text{OH})_2$ in THF gave mainly 94% II (Z = CH₂) also formed in
 similar yield from liquid COCl_2 if the resulting HCl was removed;
 III was the only by-product. The yield of III was

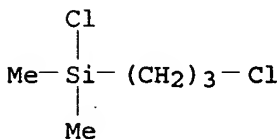
enhanced by residual HCl. I (Z = CH₂OCH₂CH₂) gave IV besides the predominantly formed disiloxane, but minor amts. of ClSiMe₂CH₂Cl, (CH₂O₂CCl)₂, ClCO₂CH₂CH₂Cl and (CH₂Cl)₂ were also found. The Si-containing phenols were inert towards COCl₂ at moderate-temps. but with added Et₃N gave HSiMe₂C₆H₄O₂CCl (m- and p-isomers). Similarly were prepared (o-ClCO₂C₆H₄SiMe₂)₂O and (o-ClCO₂C₆H₄OCH₂SiMe₂)₂O, which with the appropriate phenols and Et₃N gave (HSiMe₂C₆H₄O)₂CO (o- and p-isomers).

IT 10605-40-0P

(preparation of)

RN 10605-40-0 HCPLUS

CN Silane, chloro(3-chloropropyl)dimethyl- (6CI, 7CI, 8CI, 9CI) (CA INDEX NAME)



CC 29-6 (Organometallic and Organometalloidal Compounds)

ST phosgene silyl alc phenol addn; carbonate silyl

IT 627-11-2P 1719-57-9P 2362-10-9P 10605-40-0P

18098-85-6P 20160-66-1P 20566-53-4P 32657-05-9P

36131-27-8P 38050-04-3P 38050-06-5P 38050-07-6P

41556-35-8P 41912-68-9P 41912-71-4P 41912-74-7P

41912-76-9P

(preparation of)

L15 ANSWER 36 OF 37 HCPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1961:48296 HCPLUS

DOCUMENT NUMBER: 55:48296

ORIGINAL REFERENCE NO.: 55:9262i, 9263a-i, 9264a-d

TITLE: Addition of silicon hydrides to olefinic double bonds. V. The addition to allyl and methallyl chlorides

AUTHOR(S): Ryan, John W.; Menzie, Gerald K.; Speier, John L:

CORPORATE SOURCE: Dow Corning Corp., Midland, MI

SOURCE: Journal of the American Chemical Society (1960), 82, 3601-4

CODEN: JACSAT; ISSN: 0002-7863

DOCUMENT TYPE: Journal

LANGUAGE: Unavailable

AB cf. CA 54, 10916e. The addition of Cl₃SiH (I), MeCl₂SiH (II), and Me₂ClSiH (III) to CH₂:CHCH₂Cl (IV) and CH₂:CMeCH₂Cl (V) was studied. In the presence of H₂PtCl₆.6H₂O (VI), each hydride formed CH₂:CHMe (VII) from IV as well as Me₃-nClnSi(CH₂)₃Cl (VIII) and Me₃-nClnSiPr (IX). With V, little or no CH₂:CMe₂ (X) or isobutylylsilanes were formed, and high yields of Me₃-nClnSiCH₂CHMeCH₂Cl (XI) were obtained. No adducts isomeric with VIII or XI were detectable. The temperature had no effect on the distribution of products or yields between 40 and 140°. With VI as catalyst no reaction between VIII and the Si hydrides occurred under addition conditions. Inasmuch as PhCH₂Cl did not react with II and VI nor did V form X, VII possibly formed from IV via a 6-membered ring complex, unfavorable to V owing to orientation of hydride H to the tertiary C atom. IX may have

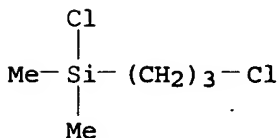
formed by addition of Si hydrides to VII. With excess IV, formation of IX was reduced, not due, apparently, to a reduction in the amount of VII formed. With 0.5% Pd on Al₂O₃, a mixture of IV and II at reflux gave VII and MeSiCl₃ almost quant. Formation of siloxanes or of alkoxy silanes from VIII and XI was straightforward. The chloroalkyl groups were not attacked by dilute acids or bases. Two methods, A and B, were used to prepare VIII and XI: Method A: A mixture of 0.5 mole IV, 0.5 mole I, and about 10-5 mole VI was refluxed; During 1 hr. the reflux temperature rose from 35 to 65° and a mixture of 1.1:1 (mole ratio) IV-I was added slowly at 75-80°, the evolved VII and volatilized reactants were trapped and the reactants returned to the flask, and the mixture was distilled to give, from 829 g. I and 421 g. IV, recovered I, 241 g. SiCl₄, b. 56°, 109 g. IX (n = 3), b. 122-3°, and 768 g. VIII (n = 3). Method B: A mixture of 1150 g. II, 637 g. IV, and 1.1 + 10-5 mole VI was pumped into a 20-ft. coil of 0.5-in. stainless steel in a bath at 125-6° and allowed to leave the coil through a spring-loaded relief valve set to maintain a pressure of 500 lb./sq. in., and at equilibrium a 429-g. sample was collected during 26 min. and analyzed by distillation to give 21 g. VII, 63 g. II, 23 g. IV, 90 g. MeSiCl₃, 54 g. IX (n = 2) (b. 125-7°, n_{25D} 1.4221), and 178 g. VIII (n = 2); at a 0.83:1 mole ratio II-IV, the yield of VIII rose to 63%, based on IV consumed. The following chlorosilanes were prepared [compound (n), method, % yield, b.p./mm., n_{25D}, d₂₅, and R_n given]: VIII (3), A, 66, 181.5°/750, 1.4638, 1.354, 0.2041; VIII (2), B, 60, 186°/750, 1.4597, 1.199, 0.2278; VIII (1), B, 50, 179°/750, 1.4488, 1.043, 0.2572; XI (3), A, 92, 194°/750, 1.4662, 1.310, 0.2112; XI (2), B, 100, 106°/40, 1.4620, 1.168, 0.2352; XI (1), B, 87, 89°/25, 1.4522, 1.030, 0.2619. Treatment of VIII (n = 2), with excess MeMgBr in Et₂O gave 85% VIII (n = 0), b. 151.5°, n_{25D} 1.4288, d₂₅ 0.8718, RD 0.2956. Similarly, XI (n = 2) yielded 71% XI (n = 0), b₇₃ 93°, n_{25D} 1.4345, d₂₅ 0.8769, RD 0.2973; the nuclear magnetic spectrum of the product indicated CH₂Cl, C-H, C-Me, -SiCH₂, and SiMe₃ groups. A mixture of 354 g. PhCl₂SiH and 200 g. IV was added slowly to 10-5 mole of VI, and the mixture left overnight and distilled to give 144 g. PhSiCl₃ (b₄₀ 105°), 31.4 g. PhCl₂SiPr (b₁₀ 105°, n_{25D} 1.5137, d₂₅ 1.125, RD 0.2669), and 313 g. PhCl₂Si(CH₂)₃Cl (b₁₀ 141-2°, n_{25D} 1.5332, d₂₅ 1.241, RD 0.2502). MeOH (550 g.) was added through a glass tube to the bottom of a flask containing 1535 g. VIII (n = 2), at 65° with stirring, the system devolatilized at 150-200 mm., and the mixture distilled through a Vigreux column to yield 1130 g. crude Me(MeO)₂Si(CH₂)₃Cl, not free of hydrolyzable Cl; 100 ml. MeOH was added, the mixture saturated with NH₃, filtered, and distilled to give pure material. Data for this compound and alkoxy silanes prepared similarly were as follows [starting compound (n), alc., % yield, b.p./mm., n_{25D}, d₂₅, RD]: VIII (3), MeOH, 97, 195°/750, 1.4183, 1.077, 0.2341; VIII (2), MeOH, 78, 185°/750, 1.4242, 1.019, 0.2505; VIII (1), MeOH, 75, 169.5°/751, 1.4283, 0.953, 0.2698; VIII (3), EtOH, 85, 124°/30, 1.4175, 1.002, 0.2512; VIII (2), EtOH, 76, 109°/30, 1.4232, 0.973, 0.2618; VIII (1), EtOH, 69, 87°/30, 1.4270, 0.932, 0.2755; XI (3), MeOH, 90, 202.5°/747, 1.4223, 1.059, 0.2401; XI (2), MeOH, 94, 193.5°/756, 1.4289, 1.009, 0.2555; XI (1), MeOH, 89, 181°/751, 1.4331, 0.948, 0.2741. Treatment of VIII (n = 3) with MeOH on a large scale yielded, in addition to 85% trimethoxy

compound, 5% $O[Si(OMe)2(CH2)3Cl]2$, b24 188°, n25D 1.4347, d25 1.136, RD 0.2293. Hydrolysis of 600 g. VIII (n = 1) with ice-water, extraction with Et2O, filtration of the dried combined extract and organic layer, and devolatilization gave 539 g. $O[SiMe2(CH2)3Cl]2$ (XII), b7 128°, n25D 1.4484, d25 0.9958, RD 0.2689. Similar hydrolysis of XI (n = 1) gave quant. $O[SiMe2CH2CHMeCH2Cl]2$ (XIII), n25D 1.4528, d25 0.9886, RD 0.2733. XIII decomposed during distillation at 13-17 mm., and after 2 hrs. at total reflux, a sample of distillate proved to be $ClMe2SiOSiMe2CH2CHMeCH2Cl$, b17 105°, n25D 1.4346, d25 1.005, RD 0.2595. Hydrolysis of VIII (n = 1), that was contaminated with IX (n = 1), yielded some $PrSiMe2OSiMe2(CH2)3Cl$, b12 104°, n25D 1.4282, d25 0.8997, RD 0.2861. Hydrolysis of 629 g. VIII (n = 2), with ice-water and extraction with C6H6 gave 407 g. $[OMeSi(CH2)3Cl]n$ (XIV), n25D 1.4709, d25 1.171, RD 0.2386, viscosity 190 cs. at 25°. Similar hydrolysis of XI (n = 2) yielded $[OMeSiCH2CHMeCH2Cl]n$, n25D 1.4700, d25 1.127, RD 0.2475. H2O (5 ml.) added dropwise to a refluxing mixture of 457 g. $(Me3Si)2O$ and 100 g. VIII (n = 1) the cooled mixture washed to pH 7 with dilute NaHCO3 and H2O, and distilled gave recovered excess $(Me3Si)2O$, 23% $Me3SiOSiMe2(CH2)3Cl$ (b39 98°, n25D 1.4189, d25 0.8991, RD 0.2808), and XII as the remainder. A mixture of 360 g. $(Me3Si)2O$, 200 g. XIV, and 10 ml. H2SO4 was refluxed overnight, cooled, washed with dilute NH4OH, dried, and distilled to give 11% $(Me3SiO)2SiMe(CH2)3Cl$ (b17 111°, n25D 1.4147, d25 0.9131, and RD 0.2741), and 7% $[Me3SiOSiMe(CH2)3Cl]2O$ (b1 115°, n25D 1.4320, d25 0.9851, RD 0.2633). To a mixture of 651 g. $Me3SiCl$ and 212 g. VIII (n = 3) was added 540 g. iso-PrOH and then 162 g. H2O, the organic layer washed with H2O to pH 7, and distilled to give 94% $(Me3SiO)3Si(CH2)3Cl$, b100 181°, n25D 1.4108, d25 0.9223, RD 0.2691. XIII (295 g.) at 260-270° for 32 hrs. liberated 10 g. volatile material, which was distilled twice and identified as X by comparison of infrared spectra and vapor pressure data. A mixture of 110 g. XIII and 0.1 g. AlCl3 at 100-135° for 30 min. gave X almost quant. A mixture of 141 g. XIII, 40 g. NaOH, 200 ml. H2O, and 200 ml. EtOH refluxed 4 hrs. yielded from a cold trap 99% methylcyclopropane, identified by the vapor pressure-temperature curve and absence of C:C in the infrared spectrum.

IT 10605-40-0, Silane, chloro(3-chloropropyl)dimethyl-
(preparation of)

RN 10605-40-0 HCPLUS

CN Silane, chloro(3-chloropropyl)dimethyl- (6CI, 7CI, 8CI, 9CI) (CA INDEX NAME)



CC 10B (Organic Chemistry: Aliphatic Compounds)

IT 98-13-5, Silane, trichlorophenyl- 141-57-1, Silane,
trichloropropyl- 594-11-6, Cyclopropane, methyl- 1628-11-1,
Silane, dichloro(3-chloro-2-methylpropyl)methyl- 2344-83-4,
Silane, (3-chloropropyl)trimethyl- 2530-87-2, Silane,
(3-chloropropyl)trimethoxy- 2550-06-3, Silane,
trichloro(3-chloropropyl)- 3401-26-1, Silane,
dichloro(3-chloropropyl)phenyl- 4518-94-9, Silane,

dichloromethylpropyl- 5089-70-3, Silane, (3-chloropropyl)triethoxy- 7787-93-1, Silane, dichloro(3-chloropropyl)methyl- 10605-40-0, Silane, chloro(3-chloropropyl)dimethyl- 13501-76-3, Silane, (3-chloropropyl)diethoxymethyl- 13508-63-9, Silane, (3-chloropropyl)ethoxydimethyl- 17256-27-8, Silane, (3-chloro-2-methylpropyl)trimethoxy- 17878-20-5, Silane, dichlorophenylpropyl- 17907-74-3, Tetrasiloxane, 3,5-bis(3-chloropropyl)-1,1,1,3,5,7,7,7-octamethyl- 17961-62-5, Disiloxane, 1-(3-chloropropyl)-1,1,3,3-tetramethyl-3-propyl- 17988-66-8, Trisiloxane, 3-(3-chloropropyl)-1,1,1,3,5,5,5-heptamethyl- 18077-31-1, Trisiloxane, 3-(3-chloropropyl)-1,1,1,5,5,5-hexamethyl-3-(trimethylsiloxy)- 18132-72-4, Disiloxane, 1,3-bis(3-chloropropyl)-1,1,3,3-tetramethyl- 18132-73-5, Disiloxane, 1,3-bis(3-chloropropyl)-1,1,3,3-tetramethoxy- 18142-53-5, Silane, trichloro(3-chloro-2-methylpropyl)- 18145-83-0, Silane, chloro(3-chloro-2-methylpropyl)dimethyl- 18171-14-7, Silane, (3-chloropropyl)methoxydimethyl- 18171-19-2, Silane, (3-chloropropyl)dimethoxymethyl- 18244-08-1, Silane, (3-chloro-2-methylpropyl)methoxydimethyl- 18244-20-7, Silane, (3-chloro-2-methylpropyl)dimethoxymethyl- 18244-32-1, Silane, (3-chloro-2-methylpropyl)trimethyl- 18291-27-5, Disiloxane, (3-chloropropyl)pentamethyl- 18388-70-0, Disiloxane, 1,3-bis(3-chloro-2-methylpropyl)-1,1,3,3-tetramethyl- 18881-68-0, Disiloxane, 1-chloro-3-(3-chloro-2-methylpropyl)-1,1,3,3-tetramethyl- (preparation of)

L15 ANSWER 37 OF 37 HCPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1954:64072 HCPLUS

DOCUMENT NUMBER: 48:64072

ORIGINAL REFERENCE NO.: 48:11303i,11304a-b

TITLE: Some new silicon compounds derived from 10-undecenoic acid

AUTHOR(S): Calas, R.; Duffaut, N.

SOURCE: Bull. mens. inform. ITERG (1953), 7, 438-40

DOCUMENT TYPE: Journal

LANGUAGE: Unavailable

AB cf. C.A. 47, 12223h. Exposure of 35 g. Me 10-undecenoate and 150 g. Et₂SiHCl 40 hrs. to ultraviolet irradiation yields 14 g. (24%) of Et₂SiCl(CH₂)₁₀CO₂Me (I), colorless, fuming liquid, b₁ 155-6°, d₂₀ 0.9666, n_{20D} 1.485; 98 hrs. irradiation increased the yield to 40%. Hydrolysis at 0° furnishes 2 parts of the corresponding silanol, Et₂Si(OH)(CH₂)₁₀CO₂Me (II), and 1 part of a siloxane, O[SiEt₂(CH₂)₁₀CO₂Me]₂ (III). II, b. 159°, d₂₀ 0.9399, n_{20D} 1.4580, viscosity at 18° 36.6 dynes/cm., III, b₁ 258-60°, d₂₀ 0.9373, n_{20D} 1.4592, saponification value 189.5, mol. weight (in camphor) 577, saponified with 0.5N alc. KOH to the corresponding diacid, b₁ 280-2°, d₂₀ 0.9566, n_{20D} 1.4701, acid number = 197. EtSiCl₂ gives by an analogous reaction after 40 hrs. of irradiation 16% (32% after 98 hrs.) EtSiCl₂(CH₂)₁₀CO₂Me, b_{0.7} 150-1°, d₂₀ 1.0429, n_{20D} 1.4611.

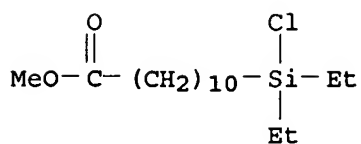
IT 18415-95-7, Undecanoic acid, 11-(chlorodiethylsilyl)-, methyl ester

(preparation of)

RN 18415-95-7 HCPLUS

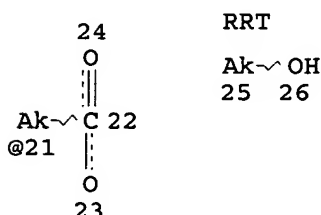
CN Undecanoic acid, 11-(chlorodiethylsilyl)-, methyl ester (8CI) (CA)

INDEX NAME)



CC 10 (Organic Chemistry)
 IT 112-38-9, 10-Undecenoic acid, silicon compounds 1825-69-0,
 Silane, chloroethoxydimethyl- 18169-88-5, Silane,
 dichloroethoxyethyl- 18171-09-0, Silane, chloroethoxydiethyl-
 18171-16-9, Silane, chlorodethoxyethyl- 18415-95-7,
 Undecanoic acid, 11-(chlorodethoxysilyl)-, methyl ester
 18416-15-4, Silanol, (10-carboxydecyl)diethyl-, methyl
 ester 18416-15-4, Undecanoic acid, 11-(diethylhydroxysilyl)-,
 methyl ester 18603-14-0, Undecanoic acid, 11-
 (dichloroethylsilyl)-, methyl ester 18765-83-8,
 13-Oxa-12,14-disilapentacosanedioic acid, 12,12,14,14-tetraethyl-
 18765-83-8, Undecanoic acid, 11,11'-(tetraethyldisiloxanylene)di-
 18765-83-8, Disiloxane, 1,3-bis(10-carboxydecyl)-1,1,3,3-
 tetraethyl- 18768-79-1, 13-Oxa-12,14-disilapentacosanedioic
 acid, 12,12,14,14-tetraethyl-, dimethyl ester 18768-79-1,
 Disiloxane, 1,3-bis(10-carboxydecyl)-1,1,3,3-tetraethyl-, dimethyl
 ester 18768-79-1, Undecanoic acid, 11,11'-
 (tetraethyldisiloxanylene)di-, dimethyl ester
 (preparation of)

=> d que 118
 L16 STR



VAR G1=X/9/16/21
 VAR G2=AK/CB
 NODE ATTRIBUTES:
 DEFAULT MLEVEL IS ATOM
 DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:
 RING(S) ARE ISOLATED OR EMBEDDED
 NUMBER OF NODES IS 25

STEREO ATTRIBUTES: NONE

L18 8 SEA FILE=CASREACT SSS FUL L16 (22 REACTIONS)

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=> d l18 1-8 bib abs fhit

L18 ANSWER 1 OF 8 CASREACT COPYRIGHT 2006 ACS on STN
 AN 141:190865 CASREACT

TI Immobilization of chiral phosphine ligands on silica gel by means of the allylsilane method and their use for catalytic asymmetric reactions

AU Aoki, Kazuko; Shimada, Toyoshi; Hayashi, Tamio

CS Department of Chemistry, Graduate School of Science, Kyoto University, Kyoto, Sakyo, 606-8502, Japan

SO Tetrahedron: Asymmetry (2004), 15(11), 1771-1777
 CODEN: TASYE3; ISSN: 0957-4166

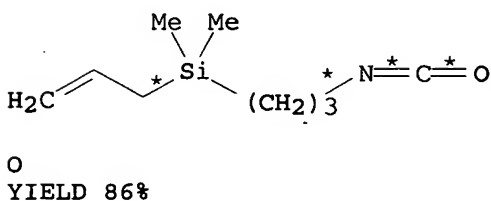
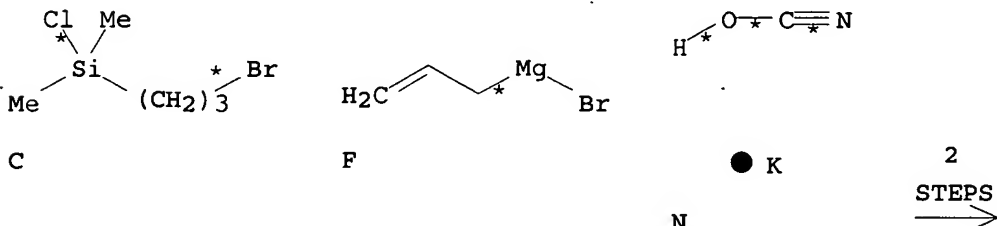
PB Elsevier Science B.V.

DT Journal

LA English

AB Three chiral phosphine ligands containing an allylsilyl group at the terminus of the side chain were prepared and immobilized on a silica gel surface by use of the allylsilane modification method. The silica-supported chiral phosphine ligands were used for rhodium-catalyzed hydrogenation and palladium-catalyzed allylic alkylation and showed high enantioselectivity.

RX(20) OF 64 COMPOSED OF RX(2), RX(4)
 RX(20) C + F + N ==> O



RX(2) RCT C 74349-12-5, F 1730-25-2

STAGE(1)

SOL 60-29-7 Et2O
 CON SUBSTAGE(1) 0 deg C
 SUBSTAGE(2) 11 hours, room temperature

STAGE(2)
 RGT H 12125-02-9 NH4Cl
 SOL 7732-18-5 Water

PRO G 738596-95-7
 NTE Grignard reaction

RX(4) RCT G 738596-95-7

STAGE(1)
 CAT 7681-11-0 KI
 SOL 68-12-2 DMF
 CON 0.5 hours, 100 deg C

STAGE(2)
 RCT N 590-28-3
 CON 0.5 hours, 100 deg C

PRO O 738596-97-9

RE.CNT 28 THERE ARE 28 CITED REFERENCES AVAILABLE FOR THIS RECORD
 ALL CITATIONS AVAILABLE IN THE RE FORMAT

L18 ANSWER 2 OF 8 CASREACT COPYRIGHT 2006 ACS on STN

AN 140:43774 CASREACT

TI Method for preparation of organodialkylalkoxysilane

IN Ramdani, Kamel; Vigin, Bernard

PA Rhodia Chimie, Fr.; Rhone Poulenc Chimie

SO Fr. Demande, 30 pp.

CODEN: FRXXBL

DT Patent

LA French

FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	FR 2841245	A1	20031226	FR 2002-7713	20020621
	FR 2841245	B1	20050218		
	FR 2841244	A1	20031226	FR 2002-15114	20021202
	WO 2004000852	A1	20031231	WO 2003-FR1921	20030623
	W:	AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW			
	RW:	GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG			
	AU 2003253076	A1	20040106	AU 2003-253076	20030623
	EP 1515977	A1	20050323	EP 2003-760774	20030623
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CN 1671719 A 20050921 CN 2003-818014 20030623
 JP 2005530855 T2 20051013 JP 2004-530906 20030623
 EP 1637534 A1 20060322 EP 2005-26550 20030623

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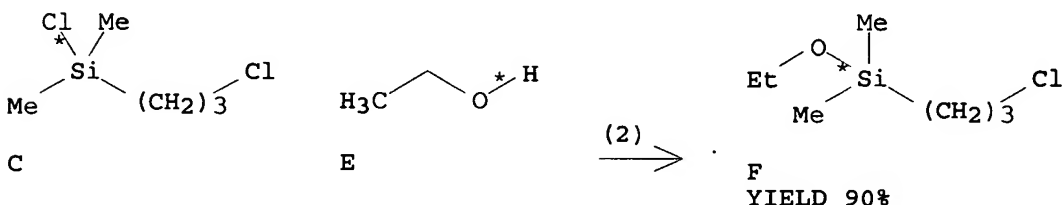
US 2005245755 A1 20051103 US 2005-518685 20050623

PRAI FR 2002-7713 20020621
 FR 2002-15114 20021202
 EP 2003-760774 20030623
 WO 2003-FR1921 20030623

OS MARPAT 140:43774

AB The preparation of organodialkylalkoxysilane is carried out by reactive distillation of an ω -haloalkyldialkylhalosilane in the presence of an alkanol. The stage of reactive distillation is implemented in a column in the presence or absence of nonreactive solvent with the removal of HCl byproduct. The ω -haloalkyldialkylalkoxysilane thus obtained is particularly useful as starting material for preparation of organosilicon compds. containing sulfur and having general formula R1OSiR2R3(CH2)3Sx(CH2)3SiR2R3OR1 by reaction of sulfurization on an alkaline metal polysulfide.

RX(2) OF 6 ...C + E ==> F...



RX(2) RCT C 10605-40-0, E 64-17-5

PRO F 13508-63-9

SOL 108-88-3 PhMe

CON 5 hours, reflux

RE.CNT 10 THERE ARE 10 CITED REFERENCES AVAILABLE FOR THIS RECORD
 ALL CITATIONS AVAILABLE IN THE RE FORMAT

L18 ANSWER 3 OF 8 CASREACT COPYRIGHT 2006 ACS on STN

AN 139.22334 CASREACT

TI Method for obtaining bis(monoorganoxysilylpropyl) polysulfides

IN Guennouni, Nathalie; Pevere, Virginie; Vigin, Bernard

PA Rhodia Chimie, Fr.

SO PCT Int. Appl., 43 pp.

CODEN: PIXXD2

DT Patent

LA French

FAN.CNT 1

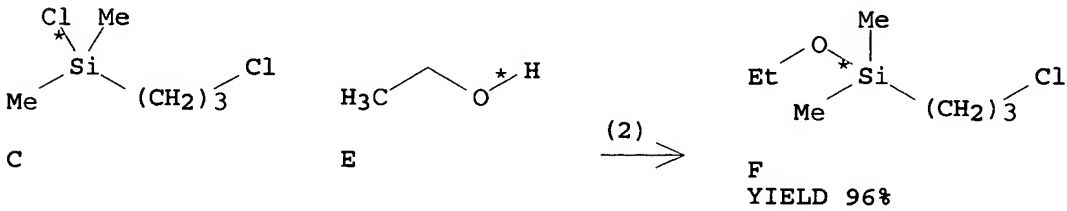
	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	WO 2003048169	A1	20030612	WO 2002-FR4204	20021206
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 SE, SG, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ,
 VC, VN, YU, ZA, ZM, ZW
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 FR 2833264 A1 20030613 FR 2001-15768 20011206
 FR 2833264 B1 20050819
 FR 2833265 A1 20030613 FR 2002-10145 20020809
 FR 2833265 B1 20060210
 AU 2002364429 A1 20030617 AU 2002-364429 20021206
 EP 1461344 A1 20040929 EP 2002-799785 20021206
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 MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ,
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 JP 2005511700 T2 20050428 JP 2003-549359 20021206
 EP 1621543 A1 20060201 EP 2005-21616 20021206
 R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE,
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 PRAI FR 2001-15768 20011206
 FR 2002-10145 20020809
 EP 2002-799785 20021206
 WO 2002-FR4204 20021206

OS MARPAT 139:22334

AB The invention concerns the preparation of bis(monoorganooxysilylpropyl) polysulfides $R_1OSiR_2R_3(CH_2)_3-Sx-(CH_2)_3SiR_2R_3OR_1$ (I, $R_1 = C_1-C_{15}$ alkyl, alkoxyalkyl; R_2 and $R_3 = C_1-C_6$ alkyl and/or phenyl; $1.5 \pm 1 \leq x \leq 5 \pm 0.1$). Said preparation is carried out by performing successively the following steps (a), (b) and (c): (a) hydrosilylation of the type: $R_2R_3HSi-Hal + CH_2:CH-CH_2-Hal \rightarrow Hal-R_2R_3Si-(CH_2)_3Hal$; (b) alcoholysis of the type: $Hal-R_2R_3Si-(CH_2)_3-Hal + R_1OH \rightarrow R_1O-R_2R_3Si-(CH_2)_3Hal$; (c) sulfidization of the type: $R_1O-R_2R_3Si-(CH_2)_3Hal + M_2Sx \rightarrow$ compound I; with Hal = halogen atom and M = alkali metal. Variations of the above reaction are also included in the invention. Thus, reaction of Me_2HSiCl with $CH_2:CHCH_2Cl$ in the presence of $[Ir(COD)Cl]_2$ (COD = 1,5-cyclooctadiene) as catalyst afforded $ClSiMe_2(CH_2)_3Cl$ (85% yield), which reacted with ethanol to give $EtOSiMe_2(CH_2)_3Cl$ (96% yield). Finally, reaction of the latter with Na_2S_4 afforded bis(monoorganooxysilylpropyl) tetrasulfide, $EtOSiMe_2(CH_2)_3-S_4-(CH_2)_3SiMe_2OEt$ (87% yield).

RX(2) OF 6 ...C + E ==> F...



RX(2) RCT C 10605-40-0

STAGE (1)

CON 150 deg C, 1 atm

STAGE (2)

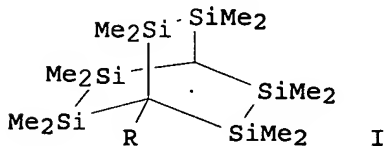
RCT E 64-17-5

CON 4.5 hours, 110 deg C, 1 atm

PRO F 13508-63-9

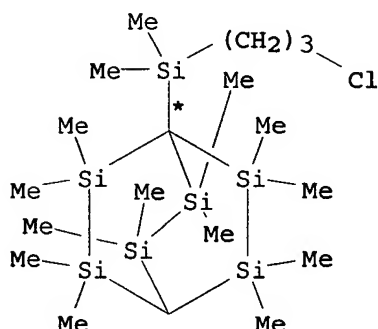
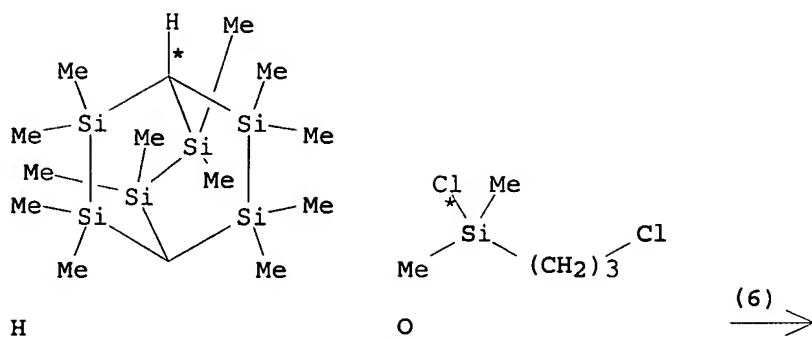
RE.CNT 10 THERE ARE 10 CITED REFERENCES AVAILABLE FOR THIS RECORD
ALL CITATIONS AVAILABLE IN THE RE FORMAT

L18 ANSWER 4 OF 8 CASREACT COPYRIGHT 2006 ACS on STN
 AN 134:56729 CASREACT
 TI Synthesis and structures of polysilacage compounds containing a silicon-silicon inter-element linkage
 AU Shimizu, Masaki; Hiyama, Tamejiro; Matsubara, Toshiaki; Yamabe, Tokio
 CS Department of Material Chemistry, Graduate School of Engineering, Kyoto University, Kyoto, 606-8501, Japan
 SO Journal of Organometallic Chemistry (2000), 611(1-2), 12-19
 CODEN: JORCAI; ISSN: 0022-328X
 PB Elsevier Science S.A.
 DT Journal
 LA English
 GI



AB To explore the possibility of three-dimensional σ -conjugation originating from Si-Si inter-element linkages, 2,2,3,3,5,5,6,6,7,7,8,8-dodecamethyl-2,3,5,6,7,8-hexasilabicyclo[2.2.2]octane (1) (shown as I, R = H) was synthesized as a model compound. The mol. structure of 1 was determined to be slightly distorted from an ideal bicyclo[2.2.2]octane skeleton by x-ray anal. Functionalization of 1 at bridgehead positions was achieved by treatment with superbase BuLi-t-BuOK followed by a reaction with an electrophile, e.g., the 1-monolithio derivative of 1 (I, R = Li) formed in situ from 1 and superbase (> 2 mol) underwent monosilylation with excess Me3SiCl in THF at -42° to give 96-98% yields of bridgehead monosilylated product (I, R = SiMe3). UV spectra of 1 and its derivs. demonstrated a bathochromic shift, particularly when dimensions of the mol. structure increased and a silyl or stannylyl group was introduced at the bridgehead. This fact was understood in terms of three-dimensional σ -conjugation between Si-Si linkages. Ab initio theor. MO calcns. of model structures of the cage compds. were also described.

RX(6) OF 70 ...H + O ==> P



P
YIELD 86%

RX(6) RCT H 218932-30-0

STAGE(1)

RGT D 109-72-8 BuLi, L 865-47-4 t-BuOK
SOL 109-99-9 THF

STAGE(2)

RCT O 10605-40-0

PRO P 313473-17-5
NTE regioselective

RE.CNT 22 THERE ARE 22 CITED REFERENCES AVAILABLE FOR THIS RECORD
ALL CITATIONS AVAILABLE IN THE RE FORMAT

L18 ANSWER 5 OF 8 CASREACT COPYRIGHT 2006 ACS on STN
AN 134:5065 CASREACT
TI The Novel Silatecan 7-tert-Butyldimethylsilyl-10-hydroxycamptothecin Displays High Lipophilicity, Improved Human Blood Stability, and Potent Anticancer Activity
AU Bom, David; Curran, Dennis P.; Kruszewski, Stefan; Zimmer, Stephen G.; Strode, J. Thompson; Kohlhagen, Glenda; Du, Wu; Chavan, Ashok J.; Fraley, Kimberly A.; Bingcang, Alex L.; Latus, Lori J.; Pommier, Yves; Burke, Thomas G.
CS Department of Chemistry, University of Pittsburgh, Pittsburgh, PA,

15260, USA

SO Journal of Medicinal Chemistry (2000), 43(21), 3970-3980
CODEN: JMCMAR; ISSN: 0022-2623

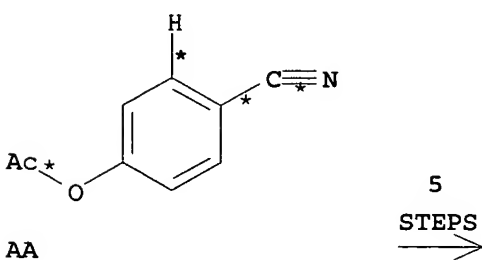
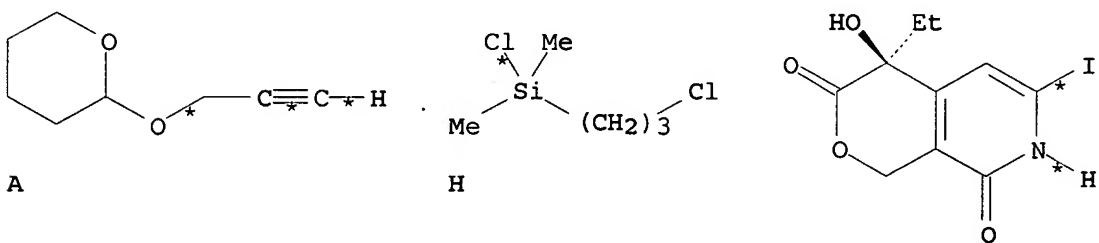
PB American Chemical Society
DT Journal

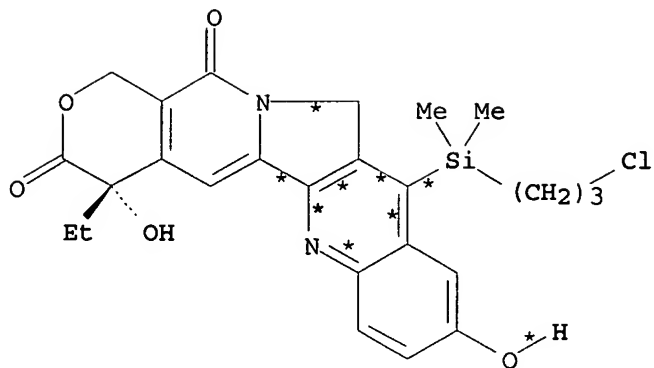
LA English

AB The rat:

The rational design and synthesis of β - and α,β -ring-modified camptothecins are described. The key α -hydroxy- δ -lactone pharmacophore in 7-tert-butyldimethylsilyl-10-hydroxycamptothecin (DB-67, I) displays superior stability in human blood when compared with clin. relevant camptothecin analogs. In human blood I displayed a $t_{1/2}$ of 130 min and a percent lactone at equilibrium value of 30%. The tert-butyldimethylsilyl group renders the new agent 25-times more lipophilic than camptothecin, and I is readily incorporated, its active lactone form, into cellular and liposomal bilayers addition, the dual 7-alkylsilyl and 10-hydroxy substitution enhances drug stability in the presence of human serum albumin. Thus, the net lipophilicity and the altered human serum albumin interactions together function to promote the enhanced blood stability. In vitro cytotoxicity assays using multiple different cell lines derived from eight distinct tumor types indicate that I is of comparable potency to camptothecin and 10-hydroxycamptothecin, as well as the FDA-approved camptothecin analogs topotecan and CPT-11. In addition, cell-free cleavage assays reveal that I is highly active and forms more stable top1 cleavage complexes than camptothecin or SN-38. The impressive blood stability and cytotoxicity profiles for I strongly suggest that I is an excellent candidate for addnl. *in vivo* pharmacol. and efficacy studies.

RX(38) OF 38 COMPOSED OF RX(2), RX(4), RX(6), RX(11), RX(12)
RX(38) A + H + O + AA ==> AJ





AJ
YIELD 44%

RX(2) RCT A 6089-04-9

STAGE(1)

RGT D 109-72-8 BuLi
SOL 109-99-9 THF

STAGE(2)

RCT H 10605-40-0

STAGE(3)

RGT E 12125-02-9 NH4Cl
SOL 7732-18-5 Water

PRO I 307925-30-0

RX(4) RCT I 307925-30-0

STAGE(1)

RGT K 603-35-0 PPh3, L 7726-95-6 Br2
SOL 75-09-2 CH2Cl2

STAGE(2)

SOL 7732-18-5 Water

PRO N 220913-64-4

RX(6) RCT O 173442-34-7

STAGE(1)

RGT Q 7646-69-7 NaH
SOL 110-71-4 (CH2OMe)2, 68-12-2 DMF

STAGE(2)

RGT V 7447-41-8 LiCl

STAGE(3)

RCT N 220913-64-4

PRO U 220913-52-0

RX(11) RCT U 220913-52-0, AA 13031-41-9
 RGT Y 661-69-8 Me3SnSnMe3
 PRO AI 307925-31-1
 SOL 71-43-2 Benzene
 NTE photochem.

RX(12) RCT AI 307925-31-1

STAGE(1)
 RGT AD 584-08-7 K2CO3
 SOL 67-56-1 MeOH, 7732-18-5 Water

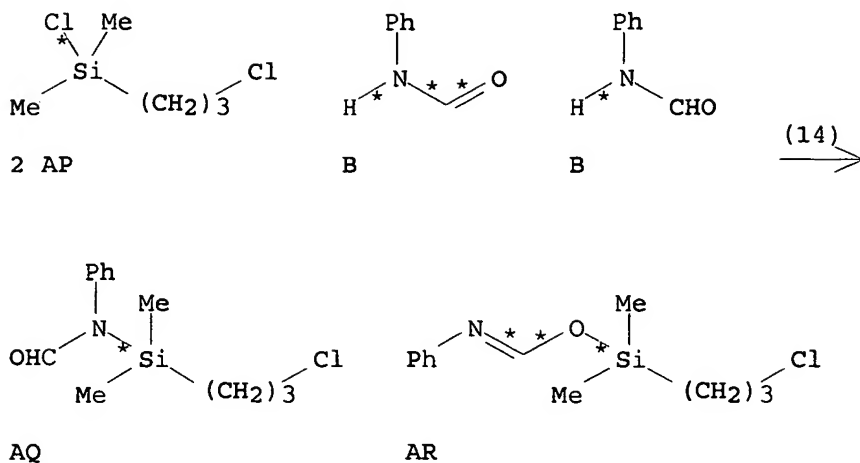
STAGE(2)
 RGT AE 64-19-7 AcOH

STAGE(3)
 RGT AF 7647-14-5 NaCl
 SOL 7732-18-5 Water

PRO AJ 302778-98-9

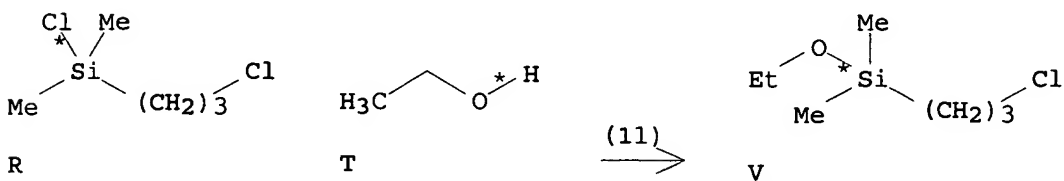
RE.CNT 34 THERE ARE 34 CITED REFERENCES AVAILABLE FOR THIS RECORD
 ALL CITATIONS AVAILABLE IN THE RE FORMAT

L18 ANSWER 6 OF 8 CASREACT COPYRIGHT 2006 ACS on STN
 AN 112:217045 CASREACT
 TI Synthesis, structure, and dynamics of (organosilyl) anilides
 AU Otter, Julie C.; Adamson, Christine L.; Yoder, Claude H.;
 Rheingold, Arnold L.
 CS Dep. Chem., Franklin and Marshall Coll., Lancaster, PA,
 17604-3003, USA
 SO Organometallics (1990), 9(5), 1557-62
 CODEN: ORGND7; ISSN: 0276-7333
 DT Journal
 LA English
 AB (Organosilyl)formanilides HCONPhSiR1R2R3 (R1R2R3 = Me2H, MePhH,
 Me3, Et3, Pr3, (i-Pr)3, Bu3, (OEt)3, (OSiMe3)3, Me2OMe, Me2Et,
 Me2CH:CH2, Me2-i-Pr, Me2C3H6Cl, Me2C2H4OAc, MeBu2, Me2Ph,
 Ph2-t-Bu) and (organosilyl)acetanilides CH3CON(p-R4C6H4)SiMe2H (R4
 = OMe, H, Cl) were prepared by amination and transsilylation. Most
 of the (organosilyl)formanilides exist as rapidly equilibrating
 mixts. of amide and imide tautomers and exhibit hindered
 rotation about the C-N bond in the amide tautomer. Bulky groups
 and alkoxy groups at silicon favor the imide tautomer. The size
 of the silyl group has no effect on the barrier to either silyl
 tautomerism or hindered rotation, while electron-withdrawing
 alkoxy groups on the silicon lower both barriers. The effect of
 substituents on the rate of tautomerism is consistent with an
 intramol., concerted mechanism. The rotamer populations are
 relatively insensitive to variations in the silyl group. The more
 stable rotamer has the silyl moiety cis to the carbonyl. The
 (dimethylsilyl)acetanilides also exist as a dynamic mixture of amide
 and imide tautomers. The attempted preparation of the SiMe2CHCl2
 formanilide derivative led to substitution at carbon rather than
 silicon. The product, (HCONPh)2CHSiMe2Cl, was shown by x-ray
 crystallog. to have distorted trigonal-bipyramidal geometry at Si,
 with two nonequivalent dative bonds from carbonyl oxygen atoms to
 silicon.



RX(14) RCT AP 10605-40-0, B 103-70-8
 RGT E 121-44-8 Et3N
 PRO AQ 126375-90-4, AR 126376-07-6
 SOL 109-99-9 THF

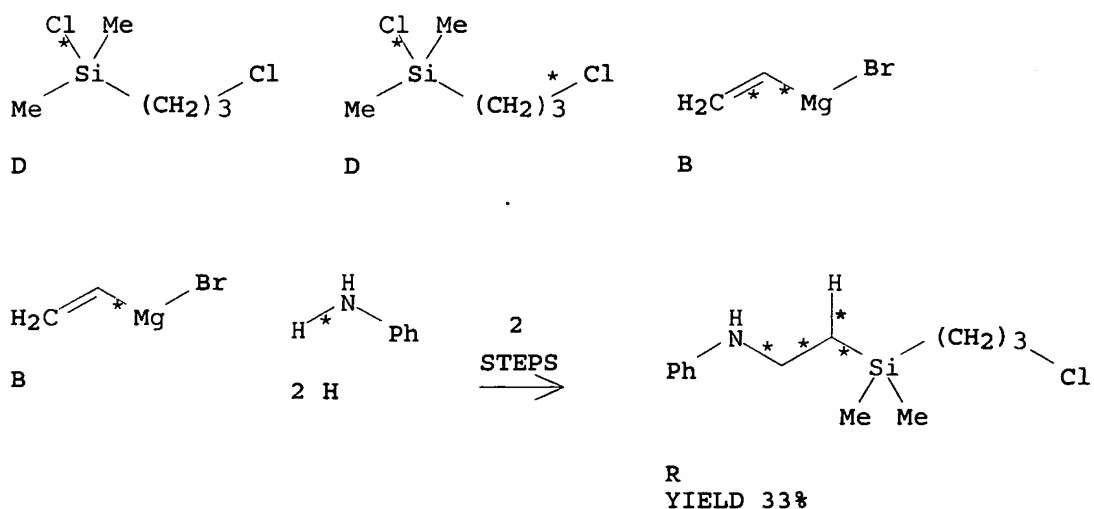
L18 ANSWER 7 OF 8 CASREACT COPYRIGHT 2006 ACS on STN
 AN 111:174221 CASREACT
 TI Synthesis of siloxyphosphines
 AU Urbaniak, W.; Marciniec, B.
 CS Fac. Chem., A. Mickiewicz Univ., Poznan, 60-780, Pol.
 SO Synthesis and Reactivity in Inorganic and Metal-Organic Chemistry
 (1988), 18(7), 695-703
 CODEN: SRIMCN; ISSN: 0094-5714
 DT Journal
 LA English
 AB Siloxyphosphines $[Ph_2P(CH_2)_nSiMe_2]_2O$ ($n = 1, 3$) were prepared by phosphidation of $X(CH_2)_nSiMe_2OEt$ ($X = Br, n = 1; X = Cl, n = 3$) with Ph₂PLi followed by hydrolysis. Free-radical addition of Ph₂PH to suitable vinylsiloxanes yielded $(Ph_2PCH_2CH_2SiMe_2)_2O$ and $(Ph_2PCH_2CH_2SiMe_2O)_2SiMeCH_2CH_2PPh_2$.

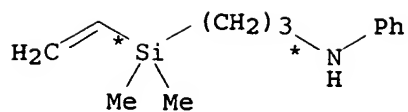


RX(11) RCT R 10605-40-0, T 64-17-5
 PRO V 13508-63-9
 NTE Petroleum ether solvent

L18 ANSWER 8 OF 8 CASREACT COPYRIGHT 2006 ACS on STN
AN 108:150547 CASREACT
TI Aminomercuration-demercuration of dimethyl(chloroalkyl)alkenylsilanes as a route to azasilacycloalkanes
AU Voronkov, M. G.; Kirpichenko, S. V.; Abrosimova, A. T.; Albanov, A. I.; Keiko, V. V.; Lavrent'ev, V. I.
CS Siberian Div., Inst. Org. Chem., Irkutsk, 664033, USSR
SO Journal of Organometallic Chemistry (1987), 326(2), 159-67
CODEN: JORCAI; ISSN: 0022-328X
DT Journal
LA English
AB 3,3-Dimethyl-1-phenyl-1-aza-3-silacyclopentane and 3,3,5-trimethyl-1-phenyl-1-aza-3-silacyclopentane were obtained by the reaction of dimethyl(chloromethyl)vinylsilane and dimethyl(chloromethyl)allylsilane with aniline in THF in the presence of mercury acetate followed by reduction with sodium borohydride. Aminomercuration-demercuration of dimethyl(3-chloropropyl)vinylsilane and dimethyl(3-chloropropyl)allylsilane results in the corresponding 3-chloropropylphenylaminoalkyl derivs. Dimethyl(3-chloropropyl)(2-phenylaminopropyl)silane undergoes cyclization under the same reaction conditions giving 2,4,4-trimethyl-1-phenyl-1-aza-4-silacycloheptane in low yield. Competitive nucleophilic substitution of the chloroalkyl group of initial silanes by aniline affords dimethyl(phenylaminoalkyl)alkenylsilanes.

RX(16) OF 22 COMPOSED OF RX(2), RX(7)
RX(16) 2 P + 2 B + 2 H ==> R + S





S
YIELD 5%

RX (2) RCT D 10605-40-0, B 1826-67-1
PRO E 88820-71-7

RX (7) RCT E 88820-71-7, H 62-53-3

STAGE (1)
RGT K 1600-27-7 Hg(OAc)₂
SOL 109-99-9 THF

STAGE (2) RGT L 1310-73-2 NaOH, N 16940-66-2 NaBH4

PRO R 113619-60-6, S 113619-61-7